Deadline for Proposals for SOT 2014 Annual Meeting Sessions: April 30, 2013

Why Submit a Proposal?
1. To present new developments in toxicology.
2. To provide attendees an opportunity to learn about state-of-the-art technology and how it applies to toxicological research.
3. To provide attendees an opportunity to learn about the emerging fields and how they apply to toxicology.

Session Types

**Continuing Education**—Emphasis on quality presentations of generally accepted, established knowledge in toxicology

*Note: CE courses will be held on Sunday.*

**Symposia**—Cutting-edge science; new areas, concepts, or data

**Workshops**—State-of-the-art knowledge in toxicology

**Roundtables**—Controversial subjects

**Historical Highlights**—Review of an historical body of science that has impacted toxicology

**Informational Sessions**—Scientific planning or membership development

**Education-Career Development Sessions**—Sessions that provide the tools and resources to toxicologists that will enhance their professional and scientific development

**Regional Interest**—Central topics of relevance that describe public health and/or ecological problems of a particular region

Submit your proposal online at www.toxicology.org
Preface

This issue is devoted to the abstracts of the presentations for the Continuing Education courses and scientific sessions of the 52nd Annual Meeting of the Society of Toxicology, held at the Henry B. Gonzalez Convention Center, March 10–14, 2013.

An alphabetical Author Index, cross referencing the corresponding abstract number(s), begins on page 536.

The issue also contains a Keyword Index (by subject or chemical) of all the presentations, beginning on page 561.

The abstracts are reproduced as accepted by the Scientific Program Committee of the Society of Toxicology and appear in numerical sequence.

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This abstract book has been produced electronically by ScholarOne, Inc. Every effort has been made to faithfully reproduce the abstracts as submitted. The author(s) of each abstract appearing in this publication is/are solely responsible for the content thereof; the publication of an article shall not constitute or be deemed to constitute any representation by the Society of Toxicology or its boards that the data presented therein are correct or are sufficient to support the conclusions reached or that the experiment design or methodology is adequate. Because of the rapid advances in the medical sciences, we recommend that independent verification of diagnoses and drug dosage be made.
The presentation will review the immunobiology of antibodies and Fc receptors and the field of therapeutic antibody development and advances in “anti-body engineering” leading to the development of improved therapeutics. A basic overview will be provided on the structure and function of antibodies, as well as the various types and formats of antibody therapeutics and technological methods of production. In addition, the immunobiology of human leukocytic Fc receptors will be discussed. These receptors serve to link humoral immune responses to cellular activities within the immune system, and generally function as either antibody-binding receptors that trigger immune cell effector functions, or as transport receptors (FcRn). Highlights will include how immunoglobulin Fc sequences are now being tailored to trigger specific Fc receptors to improve therapeutic outcomes by introducing amino acid mutations, glycoengineering, or other approaches leading to next generation formats. Known species differences in immunoglobulins and Fc receptors that may be important for pharmacologic and toxicologic evaluations will be explored, as well as other challenges in assessing the nonclinical toxicities of new antibody formats. Building upon these basic themes, the presentation will explore the current landscape of approved therapeutics and forecasts for future developments in the field. The course will provide something for those seeking basic knowledge in the field of immunology and therapeutic antibody development, as well as those seeking to refresh and enhance their knowledge of recent advances.
Toxic Effects of Metals.

M. F. Hughes¹ and M. P. Waalkes¹.¹,¹NIHEHS, Research Triangle Park, NC; ²US EPA, Research Triangle Park, NC.

Human exposures to metals are a daily occurrence because of their natural presence in the environment—their use in production of many commercial products—are byproducts of energy production and are found in many hazardous waste sites. The objective of the course is to highlight the fundamentals of metals toxicology. Metals have unique chemical and physical properties that distinguish them from organic-based chemicals. Even though some metals are essential to life, overexposure to these and other metals may result in a toxic effect in one or more organ systems. Upon exposure, metals may be absorbed, distributed throughout the systemic circulation, metabolized, and eliminated. The response of an organism following exposure to metals may be protective (e.g., induction of the metal-binding protein metallothionein), or toxicological by several mechanisms including oxidative stress. Key organ systems such as the central nervous system, the vascular system, as well as the skeleton system are affected by metals including manganese, lead, aluminum, and others. Accumulation of metals in bone has recently gained renewed interest as an eventual source of internal exposure. Noninvasive methods such as neutron activation are now being used to quantitate bone metal levels. Metals can influence gene expression, signal transduction, and epigenetics. Various toxic and carcinogenic metals such as arsenic and chromium alter the epigenetic program in cells; these effects on DNA methylation, histone tail modifications, and microRNA may be involved in metal-induced toxicity. Metals are known to cause cancer by several proposed mechanisms, including oxidative stress and the cancer stem cell hypothesis. Recent evidence suggests that developmental exposure to metals may affect stem cell population dynamics, which could result in adult onset of cancer. Overall, this is intended to be a basic course on metals toxicology, and is ideal to those who desire knowledge on the health effects of metals and useful tools used in metals toxicology research.

Advances in Nanotoxicology—Challenges.

S. M. Hussain¹ and S. F. Ali².¹US Air Force, Wright-Patterson AFB, OH; ²US FDA-NCTR, Jefferson, AR.

Recent developments in nanotechnology have generated a degree of apprehension concerning the potential risk to human health and the environment from manufactured nanomaterials (MN). The unique chemical and physical properties of MN, coupled with their high surface area per unit mass, require an extensive suite of characterization tools to effectively assess the toxicity of MN. Not only must the size and surface area of the MN be characterized prior to cellular exposure, but also a number of other specific features must be additionally evaluated, such as the size distribution, chemical composition, crystallinity, surface structure, shape, and solubility. The ionic strength of biological fluids may produce MN instability, resulting in environmental-specific aggregation tendencies that may impact toxicological results. Since aggregation of MN can modify uptake rates, transport properties, and clearance by the cell or organ system, it is critical to interpret the data from MN toxicity experiments with a detailed knowledge of the physicochemical properties of the MN at all experimental time points. Due to the lack of standardized methods to determine the physicochemical behavior of MN in biological systems, the mechanisms and nature of acute or chronic toxicity of engineered MN cannot be fully understood at this time. An understanding of a proper manner by which MN should be introduced to a biological environment has yet to be established, and consistency between cellular assay techniques has not been verified—both situations presenting clear challenges that must be addressed. This course raises issues to consider for the toxicity assessment of MN, and addresses recent advances and technical obstructions associated with conducting or interpreting in vitro or in vivo toxicity studies. The goal is to provide a comprehensive understanding of MN characterization, as well as facilitate valuable discussions of key challenges and advancements in the newly emerging field of nanotoxicology.

Gonadal Development, Function, and Toxicology.

B. McIntyre² and J. A. Flaws¹.¹NIHEHS, Research Triangle Park, NC; ²University of Illinois Urbana-Champaign, Urbana, IL.

The course objectives are to provide the basic tools for toxicologists who desire a better understanding of how to assess the effects of toxicants on the male and female gonads from development through adulthood. A focus on reproductive biology, study design considerations, reproductive endpoints, data interpretation, and use of data in risk assessment will be highlighted. Reproductive toxicity studies are among the most complex and challenging studies in the field of toxicology. The studies assess multiple interrelated endpoints of male and female reproductive development and function. To properly design, conduct, and interpret these studies, a fundamental knowledge of male and female gonadal development, anatomy, physiology, and endocrinology is required. Individual lectures will discuss the anatomy and physiology of the male and female gonads, as well as endocrine regulation of these systems. Evaluation of toxicity endpoints to assess male and female reproductive function will also be discussed, including folliculogenesis, spermatogenesis, hormone analysis, cyclicity, fertility, histopathology, and proper use of statistical analysis. The regulatory expectations related to reproductive toxicity testing, interpretation of results, and how these results are ultimately used to assess potential risks to human reproduction, will be presented. The course will conclude with methodologies for in vitro reproductive toxicity assessments for screening and investigation of mode of action. In summary, key information required for the design of reproductive toxicity studies and interpretation of reproductive toxicity data, and provide guidance for use of the data for risk assessment of reproduction, will be presented.

The Reach Regulation and Safety Assessment Approaches for Chemicals That Come in Contact with the Skin.

L. Mortensen¹ and J. Heylings².¹CIToxLAB Scantox, Lille Skensved, Denmark; ²Dermal Technology Laboratory Ltd., Keele University Science Park, United Kingdom.

REACH (Registration, Evaluation, Authorization, and Restriction of Chemical substances) is the European Union regulation on chemicals and their safe use, which came into force on June 1, 2007. The aim of REACH is to improve the protection of human health and the environment through better and earlier identification of the intrinsic properties of chemical substances. REACH places greater responsibility on the industry to manage the risks from chemicals, and to provide information on their substances. The regulation will continue fully into force in the period up to 2018. Under REACH, 30-40,000 new and existing chemicals will have to be (re)classified and registered. The regulation requires companies to conduct risk assessment and safety classification, with a minimal use of experimental animals, and to share information via databases managed by the European Chemicals Agency (ECHA). The skin (together with the respiratory system) is important as a route of chemical exposure, and as a target organ for toxicity induced by chemicals. Since under REACH so many chemicals need to be evaluated, it is important to use and develop testing methods that reliably predict human exposure and safety, while minimizing the use of experimental animals. An overview of the REACH regulation, and its practical implications for toxicological safety evaluation of chemicals marketed in Europe, will be given. Efforts to develop new methods and validation status of alternative methods that will limit the number of experimental animals to be used will be highlighted. Specifically, state-of-the-art investigational methods within dermal toxicology will be discussed since the skin is very important, both as a barrier to exposure and as a target organ. Practical examples of the use of the collected dermal safety data in the risk assessment of chemicals under REACH will be presented.

T4: Tools and Technologies in Translational Toxicology.

D. L. Mendrick² and V. S. Vaidya¹.¹Harvard Medical School, Boston, MA; ²US FDA-NCTR, Jefferson, AR.

The last decade has seen revolutionary advances in the tools and technologies available for biomedical scientists such that researchers can now conduct transformative experiments to solve unmet medical needs moving from a single cell to whole organism, and vice versa. Novel tools and innovative technologies have facilitated the development of sophisticated molecular diagnostics, enabled the use of new approaches in safety evaluation and risk assessment, and led to the development of targeted therapeutics. The development and utilization of novel technologies and tools requires interaction between scientists of differing backgrounds and talents (e.g., biologists, chemists and programmers). Only with this shared effort can medicine transform itself to meet the needs of the 21st century. The panel of experts will decode and demystify the potential of these translational and transformative technologies over a wide variety of applications, including safety/efficacy screening of compounds, imaging, omics, and in silico modeling. The key goals are to enable you to understand how recent advances in "T4": 1) help in solving important problems that have been critical barriers to progress in the field and, 2) transform the field by generating foundational resources that will be widely used throughout biomedical science for safety evaluation and risk assessment.
12 Understanding Toxic Neuropathy in Drug Development: Both Clinical and Nonclinical Perspectives.

M. Kallman1 and I. Benitez2. 1Genome Research Laboratory, Greenfield, IN; 2Vanderbilt University, Nashville, TN.

The topic of risk assessment of peripheral neuropathies is timely due to the increased clinical incidence of challenges related to multiple antecedents for the clinical presentation of neuropathies. The integration of both nonclinical and clinical dialogue on peripheral neuropathies will provide greater possibilities for successful drug development and improved patient outcomes. Peripheral nervous system toxicity is a common complication of exposure to industrial chemicals and drugs such as chemotherapeutics. Neuropathy can be caused by either limited or long-term exposure to drugs or chemicals, and toxic neuropathies can be classified by their presentation (e.g. motor vs. sensory), their electrodagnostic features or their neuroanatomical location within the peripheral nerve. Identification of toxic neuropathies prior to human exposure in the drug development process requires a multidisciplinary approach. Presentations will include information on the preclinical and clinical syndromes that have been characterized and the specific techniques for assessment. The preclinical presenters will focus on the application of preclinical data to provide risk assessment and to direct clinical assessment possibilities. The clinical presenters will emphasize the clinical situation and current treatment approaches. The course will conclude with open discussion between the presenters and the audience about opportunities for future risk assessment and the application to clinical management.


D. M. Wilson1 and A. L. Slitt. 1The Dow Chemical Company, Midland, MI; 2University of Rhode Island, Kingston, RI.

There has been an exponential increase in the attention focused on the potential role of nutrition in reducing the risk for numerous health complications, ranging from birth defects to age-associated vascular disease. Understanding the above is the increasing number of presentations and publications related to this subject, and hallmarks such as the recently revamped Food Pyramid into a Plate Icon. Chronic nutritional diseases are accepted to be a current crisis in our society; three nutrition-related diseases alone, obesity, Metabolic Syndrome, and Type 2 Diabetes, afflict over one-third of the American population. To better understand the components and etiology of nutritional diseases, it’s essential for toxicologists to be well versed in the science of nutrition. A comprehensive understanding of nutrition has broad applications in toxicology, especially considering that many of us have roles in investigating the safety of nutrients, food additives or food ingredients, studying nutritional disease, or designing and interpreting preclinical or clinical studies wherein the need to consider and understand nutritional homeostasis is essential. The potential for intersection of normal nutritional metabolic pathways with adverse outcomes is becoming even more important to delineate. This course on general nutrition, the biochemistry of nutritional pathways, the essential role of vitamins, the channeling of nutrients such as carbohydrates, proteins and fats, cellular and molecular details of nutrition, and nutritional aspects of development and reproduction, will heighten awareness of their importance in human and animal health at multiple levels. The focus will be on relevant information, starting with an introduction to nutrition, followed by a review of biochemical and metabolic reactions in nutrition, with an emphasis on their relationship to toxicology. How the nutritional status of a woman can modulate the developmental toxicity of a number of diverse toxicants, including alcohol, will be presented.

14 Genetic and Epigenetic Determinants of Susceptibility to Environmental and Occupational Toxics.

V. L. Johnson1 and B. Yuceloy2. 1BRT-Burlington Research Technologies, Morrisville, NC; 2Toxicology and Molecular Biology Branch, NIOSH/CDCC, Morgantown, WV.

The most common chronic disorders are multifactorial in nature, influenced by complex interactions of genetic and gene-environment interactions. While gene expression is a dynamic process that varies in response to a myriad of internal and external triggers and the surrounding microenvironment, the epigenetic mechanisms play a key role in mediating environmental influences on gene expression and epistatic interactions. In this respect, the expression of complex phenotypes should be assessed in a functional context that would look at the interplay between environmental, genetic, and epigenetic factors. Recent advances in genetic and epigenetic research offer new opportunities to integrate experimental approaches, including animal models and in vitro research, with computational strategies to predict such interactions at multiple levels of complexity. The focus of this session will be on current research investigating the role of genetic factors, epigenetic factors, and gene-environment interactions in the development and outcomes of complex diseases caused by environmental and occupational toxins.

15 Genetic Susceptibility to Occupational and Environmental Exposures.

D. C. Christiani. School of Public Health, Harvard Medical School, Boston, MA.

Due to their high prevalence in the general population, genetic polymorphisms in the susceptibility genes may predispose community members exposed to toxicants. Studies in genetic susceptibilities can eventually provide the following benefits: (1) to provide mechanistic insight of the etiology of disease; (2) to identify the more susceptible subpopulations with respect to exposure; (3) to provide valuable input in setting exposure limits by taking into account individual susceptibility. Research in this area has provided promising insights to occupational medicine, such as those illustrated by the NAT2 polymorphisms-aniline dyes and bladder cancer, and the HLA-DPB1*0401 in chronic beryllium disease, a hypersensitivity-mediated inflammatory disorder. Nonetheless, even for the genetic susceptibility markers that have been shown scientifically to have a clear role in disease risk, the value of wide-scale genetic screening in occupational settings remains limited. In the general environmental setting, there are limited, but growing data on the role of common gene polymorphisms in predisposing children and adults to inflammation-related respiratory disorders induced by air pollution, and to heavy metal toxicity. The purpose of this presentation is to discuss state of knowledge with regard to gene variants interacting with environmental exposures in causing cancer and inflammatory disorders.

16 Toxicogenomic and Systems Biology Approaches in the Understanding of Toxicity and Leukemogenesis Induced by Benzene.

C. McKhail1, L. Zhang1, Q. Lan2, R. Thomas1, A. E. Hubbard1, R. Vermeulen1, G. Li1, S. M. Rappaport1, S. Yin1, M. T. Smith1 and N. Rothman1. 1School of Public Health, University of California Berkeley, Berkeley, CA; 2Division of Cancer Epidemiology and Genetics, National Cancer Institute, Bethesda, MD; 3Institute of Risk Assessment Science, Utrecht University, Utrecht, Netherlands; 4Institute of Occupational Health and Poison Control, Chinese Center for Disease Control and Prevention, Beijing, China.

Benzene is an established cause of acute myeloid leukemia (AML) and may cause one or more lymphoid malignancies in humans. Occupational exposure to benzene, even at levels below the current U.S. occupational standard of 1 ppm, causes hematotoxicity. Toxicogenomics (e.g. genomics, transcriptomics and epigenomics) and systems biology (study of the interactions among toxicogenomic endpoints using bioinformatics) approaches in human populations, animals, and in vitro models, exposed to a range of benzene levels, are key to understanding gene-environment interactions in benzene toxicity and can identify biomarkers of exposure, early effect and susceptibility. Through analysis of the peripheral blood mononuclear cell (PBMC) transcriptomes of 125 workers exposed to a wide range of benzene levels, we recently reported highly significant widespread perturbation of gene expression at all exposure levels, as well as alterations in AML and immune response pathways. Sequencing the PBMC transcriptomes from a subset of the study subjects revealed additional alterations in gene expression. From preliminary epigenomic data in the human subjects, we have identified benzene-induced alterations in the DNA methylome and miRNome. Using genomic screens in yeast, with subsequent confirmation in human cells, we have identified potential biomarkers of susceptibility. We are developing bioinformatic methods to integrate these and future toxicogenomic datasets, in a systems biology approach, to further understand pathways of benzene toxicity and to reveal potential biomarkers associated with a range of exposures.

Supported by NIH grant P42ES04705.

17 Integrated Genetic and Genomic Approaches to Understand Susceptibility to Toxicant-Induced Lung Disease.

S. R. Kleeberger. Laboratory of Respiratory Biology, NIEHS, Research Triangle Park, NC. Sponsor: V. Johnson.

Genetic background has an important role in susceptibility to complex lung diseases, and the genetic contribution to disease phenotypes varies between populations. Understanding the mechanisms of interactions between genetic background
and exposures to environmental stimuli are critical to disease prevention. Animal models, particularly inbred mice, provide important insight to understand disease etiologies because genetic background and environmental exposures can be controlled. Tools including in silico haplotypeing, collaborative cross and diversity outcross mouse panels, bioinformatic applications, and -omics technologies have enhanced our ability to identify disease genes and pathways to guide translational investigations that apply these discoveries to human populations. Epigenetic and genomic approaches have yielded important insight to mechanisms of susceptibility to many complex traits and diseases. We have integrated inbred mouse and cell-based models with haplotype association mapping (genetic), global gene expression analyses (genomic), and expression quantitative trait locus mapping (eQTL or genetical genomics) to identify candidate susceptibility genes and associated gene networks important in toxicant-induced lung injury. The overarching goal of these investigations is to determine whether human homologues of these susceptibility genes associate with disease risk in human populations. Efforts to identify and validate susceptibility genes in mouse models of environmental disease with a goal toward translational application have enabled identification of individuals who are susceptible to disease. For example, epidemiological and clinical investigations have associated functional polymorphisms in human Nrf2 (NF-E2 related factor 2) and TNF (tumor necrosis factor alpha) with susceptibility to acute lung injury and ozone-induced changes in lung function, respectively. Importantly, these discoveries may also lead to novel intervention or therapeutic strategies to prevent disease.

18 Developmental Exposure to Bisphenol A and Lead: Effects on Metabolic Homeostasis and the Epigenome.
D. Dolinoy, University of Michigan, Ann Arbor, MI.

Environmental exposures during early development and other critical life stages may induce changes to the epigenome resulting in potentially deleterious phenotypic effects including metabolic disease, cancer, and neurological disorders. The field of epigenetics is experiencing a rapid advancement in technology, methodology, and data acquisition that now allows for the identification of the constellation of genomic loci with altered epigenetic status following dose-dependent exposures. Thus, epigenomic profiling facilitates the identification of biomarkers of exposure, enabling clinicians to identify at-risk individuals prior to disease onset. Utilizing a multi-pronged approach with an in vivo mouse model, human clinical samples, and an ongoing 15-year longitudinal epidemiological study, the overall goal of this presentation is to elucidate the impact of perinatal bisphenol A (BPA) and lead (Pb) exposure on metabolic homeostasis and DNA methylation, and the interplay between the two. Developmental exposure to environmentally relevant levels of BPA has been shown to affect both global and gene-specific DNA methylation patterns in rodents. We now draw upon data from multiple whole-epigenome platforms to show that multiple dose levels of BPA affect DNA methylation in mice and humans and that these epigenetic effects are non-monotonic in dose response. Preliminary studies also indicate that Pb exhibits epigenetic effects that may contribute to its known neurotoxic and obesogenic activities.

19 Predictive Toxicology Paradigms for Understanding Carbon Nanotube Toxicity in the Lung.
J. C. Bonnet1, A. Ne1, D. W. Porter1, V. Castranova2 and K. E. Pinkerton3, 1NIOSH, Morgantown, WV; 2Department of Mechanical & Aerospace Engineering, West Virginia University, Morgantown, WV; 3Research Center for Exotic Nanocarbons, Shinshu University, Nagano, Japan.

Nanotechnology is rapidly developing, resulting in the production of a variety of engineered nanoparticles. Carbon nanotubes (CNTs) represent an important family of nanoparticles because they have many potential uses in engineering, electronics, and medicine due to their ease of functionalization, unusual strength, and electrical conductivity. However, these novel nanostructures also represent a potential human health risk, due to the possibility of inhalation exposure and evidence that the lung and cardiovascular systems are targets for hazardous effects. Inhalation studies in rodents show that CNTs deposit within the distal regions in the lungs and migrate to the pleura to cause inflammatory and/or fibrotic effects. Presentations in this session are aimed at elucidating the pulmonary and cardiovascular effects of CNTs, and how an increasing variety of functionalized CNTs can be evaluated via high-content screening. Because the toxicological activity, we will address high-content screening for the development of structure-activity relationships relevant to inhalation toxicity and safer design of nanoparticles. This will include exploration of factors that mediate toxic effects such as high aspect ratio, durability, and residual metal content and discuss how removing metal catalysts or changing surface properties alters the pattern and timing of toxicity. While the lung is a major target organ, another goal is to determine the potential for inhaled CNTs to have toxic effects that reach beyond the lung to influence the cardiovascular system. Finally, we will discuss how susceptibility factors, both genetic and environmental, determine pulmonary and cardiovascular toxicity to CNTs. The outcome of this session is to gain a better understanding of the structure-activity relationships, target organs, and susceptibility factors that will aid the development of predictive toxicology paradigms for understanding CNT toxicity.

20 Time Course of Pulmonary Responses to Inhaled Multiwalled Carbon Nanotubes.
D. W. Porter1, A. F. Hubbs2, R. R. Mercer3, N. Wi4, W. McKinney5, B. T. Chen6, M. G. Wolfarth7, L. A. Battelli8, J. F. Scabilloni9, D. Schwegler-Berry9, S. Friend3, S. Tsuuroka3, M. Endo4, D. Frazer1 and V. Castranova1, 1NIOSH, Morgantown, WV; 2Department of Mechanical & Aerospace Engineering, West Virginia University, Morgantown, WV; 3Research Center for Exotic Nanocarbons, Shinshu University, Nagano, Japan.

In the present study, an aerosol of multi-walled carbon nanotubes (MWCNT) was produced with an acoustic generator, and airborne concentration and size distribution was determined. Mice were exposed by whole body inhalation to MWCNT (5 mg/m3, 5 hours/day, 12 days) and pulmonary responses were monitored at 1 day, 2, 4, 12, 24 and 48 weeks post-exposure. Pulmonary responses were investigated using whole lung lavage, histopathology, morphometry, and enhanced darkfield light microscopy studies. MWCNT lung burden was also measured to assess MWCNT clearance. Data indicate that the lung burdens of MWCNT in this study represent lung burdens relevant to estimated human occupational exposures and caused time-dependent pulmonary inflammation, damage and pulmonary fibrosis. Using enhanced darkfield microscopy, MWCNT fibers were found in lavage of the pleural space, parietal pleura, and respiratory muscles of the diaphragm and chest wall. The time course of pulmonary responses and their relationship to MWCNT lung burden and clearance will be discussed.

21 Establishment of Carbon Nanotube Structure-Activity Relationships (SARs) That Can Be Used to Understand Pulmonary Toxicity and Safer Design.
A. Nel, UCLA, Los Angeles, CA.

There is a fundamental gap in understanding how the physicochemical properties of carbon nanotubes (CNTs) contribute to hazard generation in the lung. Without this knowledge, it is difficult to evaluate CNT safety in a predictive manner. Our goal is to develop a predictive toxicological paradigm for CNT safety assessment in which we define the structure-activity relationships (SARS) leading to hazard generation at the nano/bio interface, including ways to design safer materials that do not induce chronic inflammation and fibrosis. To achieve this goal, we are developing a series of single-wall and multi-wall CNT test materials that can be screened by robust cellular assays to perform hazard ranking and SAR analysis. We are looking at the role of CNT dimensions (including length, diameter and aspect ratio), dispersability, catalytic surface chemistry, electronic properties and purity in initiating cooperative cellular interactions in macrophages and cells of the epithelial-mesenchymal transition (EMT), which are involved in the pathogenesis of pulmonary inflammation and fibrosis. The above physicochemical characteristics impact the lysosomal stability in macrophages in a hierarchical fashion, leading to cathepsin B release and assembly of the subunits of the NALP3 inflammasome. This leads to IL-1 beta release, which primes the EMT unit and initiates a march of events leading to TGF-beta and PDGF production and subsequent induction of chronic inflammation and fibrosis in the lung. Utilizing myeloid and endothelial cell lines, it is possible to study the induction of these biomarkers in relation to the property variations of the CNT materials, predicting the SARs that are associated with pulmonary inflammation and fibrosis. Moreover, we have also implemented surface coating and functionalization approaches that can change the hazardous characteristics, leading to the design of safer CNTs. The overall utility of this research exploration is to establish a predictive and quantitative toxicological paradigm for the safety assessment of CNTs and their safe implementation in the marketplace.

22 Surface and Chemical Modification of Single-Walled Carbon Nanotubes Does Not Necessarily Create a Safer Nanomaterial.
K. E. Pinkerton, University of California Davis, Davis, CA.

We hypothesized iron (Fe) content and morphology of inhaled single-walled carbon nanotubes (SWCNTs) would influence the extent of cellular injury and alters homeostasis in the lung. Rats (SD) were exposed (1 mg/m3) to either aerosolized
SWCNTs (raw FeSWCNT or purified cSWCNT), carbon black (CB), crocidolite asbestos, or fresh air via nose-only inhalation for 6 h/4 d. SWCNTs containing varied amounts of iron. Depending on the endpoint of interest, pulmonary responses of SWCNTs sometimes followed that of CB while in other circumstances matched that of crocidolite. Notably, animals exposed to FeSWCNTs were unable to respond to an additional oxidant challenge and cSWCNTs exposed animals had a latent and persistent development of mucus cells in the distal airways. In summary, while some toxicity endpoints follow patterns comparable to CB or crocidolite, the respiratory effects of inhaled FeSWCNTs and cSWCNTs appear to be unique. These changes could be suggestive of precursor events to pathologic changes that could develop under more prolonged exposure conditions.

Pulmonary exposure to various nanoparticles has been reported to cause lung inflammation and in some cases fibrosis. This presentation describes cardiovascular responses which occur following inhalation of titanium dioxide nanospheres or multi-walled carbon nanotubes in rats. Pulmonary exposure to nano titanium dioxide inhibits the ability of systemic and coronary arterioles to respond normally to oxidant challenge and cSWCNTs exposed animals had a latent and persistent development of mucus cells in the distal airways. In summary, while some toxicity endpoints follow patterns comparable to CB or crocidolite, the respiratory effects of inhaled FeSWCNTs and cSWCNTs appear to be unique. These changes could be suggestive of precursor events to pathologic changes that could develop under more prolonged exposure conditions.

The dysfunction occurs 24 hours after exposure and declines over several days. Inhalation of multi-walled carbon nanotubes in rats. Pulmonary exposure to nano titanium dioxide inhibits the ability of systemic and coronary arterioles to respond normally to oxidant challenge and cSWCNTs exposed animals had a latent and persistent development of mucus cells in the distal airways. In summary, while some toxicity endpoints follow patterns comparable to CB or crocidolite, the respiratory effects of inhaled FeSWCNTs and cSWCNTs appear to be unique. These changes could be suggestive of precursor events to pathologic changes that could develop under more prolonged exposure conditions.

V. Castranova, Pathology & Physiology Research Branch, NIOSH, Morgantown, WV.

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Within the arteriolar smooth muscle partially reverses these effects. Inhalation of multi-walled carbon nanotubes also inhibits coronary arterial responsiveness to dilators. This dysfunction occurs 24 hours after exposure and declines over several days thereafter. Human relevance of these rat data will be discussed.

J. V. Boekelheide, Pathology & Physiology Research Branch, NIOSH, Morgantown, WV.

While studies with rodents suggest that inhaled carbon nanotubes could represent a human health risk for the development of pulmonary diseases, it is likely that individuals with pre-existing disease or those with specific genetic deficiencies would be most susceptible to exposure and lung injury. The goal of this presentation is to emphasize genetic and environmental factors that determine susceptibility to inflammatory, immune, and fibroproliferative responses in the lungs of mice exposed to carbon nanotubes. Evidence will be presented showing that pre-existing non-allergic lung inflammation induced by bacterial lipopolysaccharide (LPS) or allergic airway disease induced by ovalbumin sensitization alters the immune response and fibroproliferative effects of carbon nanotubes as well as modulates other aspects of airway remodeling such as inflammatory cell infiltration and mucus hypersecretion. In addition, transgenic mouse studies show that susceptibility or resistance to carbon nanotubes is determined by the relative abundance or expression of specific genes that encode transcription factors involved in the innate immune response. Finally, the consequence of enhancing the physical or chemical properties of carbon nanotubes by atomic layer deposition will be addressed to better understand how post-synthesis engineering affects toxicity and disease progression.

J. C. Bonner, Toxicology, NC State University, Raleigh, NC.

Translatable Indicators of Testicular Toxicity: Inhibin B, microRNAs, and Sperm Signatures.

K. Boekelheide, B. McIntyre, M. Coulson and R. E. Chapin. Brown University, Providence, RI; Pfizer Drug Safety, DART Group, Groton, CT; National Toxicology Program, NIEHS, NIH, Research Triangle Park, NC; AstaZeneca R&D, Alderley Park, United Kingdom.

The typical endpoints used in preclinical animal models for reproductive toxicity testing, such as histopathology, are not translatable for human clinical assessment, which typically focuses on the analyses of semen and serum hormones. Therefore, when testicular toxicity arises in preclinical toxicity testing, the methods currently available to monitor this liability in clinical trials are limited. Of these limiting events is a need to develop sensitive and translatable indicators that reliably reflect testicular function. This presentation describes the feasibility testing, preclinical tests of male reproductive toxicity, and current methods for assessing testicular function in men in clinical trials. The following three talks will discuss currently active efforts to develop improved translatable indicators of testicular toxicity and their potential for recovery/reversibility, and the margin of safety between the preclinical toxic exposure and the expected clinical exposure. Noninvasive (and often insensitive) biomarkers are often benchmarked against these exposures. This presentation will review the benefits of this sometimes challenging and inferential paradigm, but will also underscore that new tools and methodologies would be of value in understanding the human relevance of drug-induced testicular findings observed in animals.
been working to define which miRNAs are testis-specific or enriched, and to de-
fine their changes in the blood during testicular toxicant-induced injury. This pres-
entation will briefly review the underlying biology and present an update of the
work in this fast-moving area.

K. Boekelheide, Brown University, Providence, RI.

As a pure population of cells that have matured and differentiated within the semi-
niferous epithelium, sperm carry all of the information needed to fertilize an oocyte
and initiate embryogenesis. Sperm deliver small and large RNAs and DNA to the
oocyte. Using high throughput array technology, a panel of altered sperm miRNAs
has been identified after exposure of rats to testicular toxicants. In men, the extent
of sperm DNA methylation is related to sperm motility. The ultimate goal is to
identify translatable sperm molecular signatures that will allow the rapid assessment
of sperm effects in preclinical test species and exposed men.

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In the risk assessment process, DNA-reactive agents are generally considered to
have no thresholds for their biological effects and this assumption formed the basis
for linear low-dose extrapolation of any carcinogenic effects induced by these
agents. On the other hand, cells have evolved to handle DNA lesions from endoge-
nous and many exogenous DNA-reactive agents. In fact, DNA-repair processes are
strictly conserved from bacteria to humans underlying their importance in protec-
tion against the effects of these agents, such as disease and aging. In recent years,
low-dose response for DNA-reactive agents has been an active domain for research
in toxicology, with publication of several datasets with large numbers of doses fo-
cused on the determination of no-observed-genotoxic-effect-level (NOGEL) val-
ues. This effort has included dose-response modelling to identify the best fit be-
tween linear and nonlinear models, such as bilinear (hockey-stick) and the benchmark
dose (BMD) suite of models, and most datasets have supported a bilin-
ear or nonlinear/threshold dose response as providing best fit based on statistical
criteria. However, empirical demonstration of statistically-supported nonlinear/threshold
dose response alone is not sufficient to achieve a paradigm shift in risk assessment. A clear understanding of the biological processes behind the
shape of the low-dose response curve is similarly a critical piece of this journey, and
one where less effort has been focused to date within toxicology. This workshop will
explore some of the questions that need to be addressed to understand and bridge
DNA-repair processes and cellular responses with the mode-of-action driving these
non-linear/threshold dose-responses for genotoxic effects. The workshop will exam-
in these experimental technologies generated new biological insights. HR has emerged as an important driver of carcinogenic sequence rearrangements.
HR repairs double strand breaks in the S/G2 phases of the cell cycle resulting from
endogenous and exogenous processes. While usually homologous recombination may
result in sequence rearrangements and loss of heterozygos-
yty, both of which are prominent features of cancer cells. BER is also a key DNA re-
pair pathway. In this case, damaged bases are removed, the DNA backbone is
cleaved, the ends are processed and the resulting gap is filled from the opposite
strand. Both pathways are usually error-free, but there are still events where
there are misalignments during HR, and misinsertions during BER. Also, both
pathways are active in response to spontaneous DNA damage, and thus would be
expected to be active in response to low dose radiation.

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Bronx, NY; Sponsor: J. Klapacz.

The DNA mismatch repair (MMR) system is essential for maintaining the integ-
rity of mammalian genomes by removing misincorporated nucleotides that result from
erroneous replication. In addition, MMR participates in the early steps of check-
point activation and apoptosis during the cellular response to alkylation induced
O6MeG:T mismatches and many other DNA lesions, particularly relevant at low-
dose, clinically-relevant exposures. Mutations in mammalian MMR genes result in
increased spontaneous mutation rates and strong predisposition to colorectal can-
cers and other cancers. Eukaryotic MMR is a complex system that requires the in-
teraction of several MutS and MutL proteins for the initiation of the repair reac-
tion. Subsequent to mismatch recognition, downstream events are activated that
lead to the excision of misincorporated or damaged nucleotides and the signaling of
DNA damage-induced cell cycle arrest and apoptosis. The loss of the MMR-de-
pendent DNA damage response is of significant clinical relevance as it results in
increased resistance to many alkylating and chemotherapeutic agents. Our research
program focuses on elucidating the functions of the individual MutS and MutL ho-
nologs in mammalian MMR and on assessing their importance for tumor suppress-
sion and the DNA damage response. Our results show that the DNA damage sig-
naling by MMR is important for the suppression of tumorigenesis in the initial
stages of the process. In addition, we found that MMR mismatch mutations can ef-
cfectively separate the DNA repair and damage response functions and result in
more heterogeneous cancer phenotypes than those caused by complete loss of func-
tion mutations.
Homeostasis with cellular stress response pathways involves negative feedback acting through a series of steps to reduce stressor concentrations. Many stress response pathways have rapid response, post-translational signaling and slower signaling through transcriptional upregulation of gene families. Our laboratory, in close collaboration with scientists from Unilever Safety and Environment Assurance Centre (SEAC), UK, has examined multiple biological read-outs in cells treated with several DNA-damaging compounds in order to create mechanistic computational models for micronuclei (MN) formation across wide dose ranges. The readouts in several cell types—AH1–1, HT-1080 and TK6—include dose and time-dependent examination of whole genome gene expression, high content imaging of DNA-damage markers (H2AX and p53), semiquantitative measures of key phosphoproteins (ATM, ATR, p38, Chk2, and p53), and MN as a measure of DNA-damage. Transcriptional upregulation only occurred in the regions of dose-response with clear increases in MN formation. Post-translational activation of rate constants for DNA-repair processes acting through activation of specific kinases appears to be the main contributor to regulation of DNA-damage at lower doses of these compounds. This talk describes stress pathway homeostatic feedback loops, shows our steps to populate high- and low-dose models for DNA-damage response pathways with endosome, quercetin and methylmethanesulfonate, notes our progress in creating mechanistically-based low dose threshold models and discusses the value of these computational models in guiding new experimental approaches for assessing thresholds.

Do Alkylating Agents Cause Genotoxic Thresholds through a DNA Repair Mode of Action?

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The long-standing theory that linear dose responses exist for all DNA reactive genotoxic agents has recently been challenged. This paradigm shift towards accepting thresholds has been initiated by the scientific and regulatory community. High power non-linear dose responses are being produced in robust test systems. However, for the scientific community to accept a range of doses as biologically irrelevant, a plausible mechanism of action must be shown experimentally. Many different research groups are tackling this issue for methyl- and ethyl-methane sulphonate (EMS and MMS) and methyl- and ethyl-nitrosourea (MNU and ENU) which have all been shown to exhibit non-linear dose responses for both mutagenicity and clastogenicity in vitro and in vivo. These alkylating agents induce specific DNA adducts (O6-alkylG, N7-alkyl-G and N3-alkyl-A), and recent work has been to investigate the roles of DNA repair in relation to their genotoxic thresholds. Specific DNA repair enzymes (N-methyl DNA purine glycosylase [MPG] and O6-alkyl guanine transferase [AGT]) have been shown to be up-regulated by low dose alkylating agents, and knocking down these specific DNA repair enzymes in vitro alters the shape of the dose response e.g. to EMS and MNU. Therefore, mono-functional alkylating agents have threshold dose responses through a DNA repair mode of action.

Incorporation of Exposure Data and Chemical Properties into Early In Vitro Screening Studies: Putting Early Hazard Identification into Appropriate Context.

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Over the past several years, a multiplicity of innovative early-screening assays have been adopted to improve our ability to select drug candidates with the maximal opportunity to eventually succeed in later development. Examples of medium and high-throughput assays that can enable early hazard identification include ion channels related to cardiovascular safety (hERG, NaV1.5, and others), assays to understand CNS permeability and hepatobiliary transport (Pgp, BSEP, etc.), and receptor and kinase screens, as well as screening ADMET assays. These assays are typically employed early in the drug discovery process when little other data are available to help contextualize the early-screening result. Data output from these early screens is often produced, then utilized by project teams to select molecules for advancement, de-prioritization, or to accumulate the knowledge base around a compound. In this model, the early-screening data can be used as a starting point, such as an IC50 value, which is then stored in a database often without further consideration or re-evaluation. There is a possibility that early-screening data in such a model is underutilized or potentially misleading. An inherent problem with generating screening data is that it is often produced in the absence of accompanying data (e.g., in vivo exposure), which is information that can be leveraged to place these data into better context and enable more accurate predictions. In this workshop, several use case examples will be presented where in vivo exposure information and chemical compound properties have been woven into early in vitro toxicity screening data enabling the discovery toxicologist to conduct a more robust assessment of the hazard identification assay data.

Inclusion of Exposure Data to Early Toxicity Screening: Using the Bile Salt Export Pump (BSEP) Screen As an Illustrative Example.


Inhibition of the bile salt export pump (BSEP) is a recognized risk factor in the development of drug induced hepatic toxicity. To date there has not been an animal model that can be reliably used to predict toxicity arising from inhibition of bile salt transport. Thus, in order to provide project teams with an early hazard identification tool for BSEP inhibition, a vesicle-based BSEP assay was developed. Guidance for decision making or prioritization was initially based on a series of cutoff values, where potency on BSEP was used as a singular criterion for hazard flagging. As part of an effort to refine how early screening data are used, BSEP inhibition data was obtained from a compendium of approved and withdrawn drugs. The available human exposure data was then applied in a ratio calculation (BSEP IC50/Exposure) in order to provide an estimate of the safety multiple. Using the ratio-based data, a more robust evaluation of the IC50 data is now possible and the decision making power is enhanced. In addition, both false positive and false negative compounds were further analyzed and hypotheses generated to refine judgment around the hazard for individual molecules.

High-Content Mechanistic Screening (HCMS) Technology to Impact Safety in the Context of Efficacy, Exposure, and Toxicity.

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High content mechanism screening (HCMS) is a powerful platform that images numerous mechanistic toxicity endpoints in full dose response at various time points in cells of choice to generate IC50 based toxicity profiles presented as heatmaps. This allows to compare toxicity profiles of compounds across chemical series or within a series, with the goal to only take compounds forward that meet an acceptable safety profile and minimize the probability of attrition in later stages of development. We have successfully applied this approach to retrospective analysis for organ toxicity prediction as well as within our own portfolio. This work demonstrated that accurate prediction of clinical safety outcome depended on the availability of accurate information on clinical exposure levels. Rank ordering of compounds should be done in the context of exposure to most accurately define the safety window. When adjusted for clinical Cmax, retrospectively, the rank ordering of compounds fell in line with the FDA status and known clinical adverse events observations. This information was available to us for retrospective validation of the HCMS technology, but is not available for discovery compounds. To address this problem, as part of retrospective validation, we used industry-standard approach of using pharmacologically-based pharmacokinetic (PBPK) model and predicted human Cmax to normalize HCMS toxicity data. We demonstrated that when HCMS data were adjusted for predicted Cmax, that was calculated using PBPK model CloePK, rank order of the compounds fell in the same range as when we used clinical Cmax. In order to generate PBPK predictions a set of standard ADME data is used as an input.

Lead Optimization Against Toxicological End Points in Drug Discovery: Recognition of Structural Determinants of Small Molecule Target Organ Exposure and Toxicity.

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Target organ toxicity is a leading cause of attrition for novel small molecules in non-clinical drug discovery and early clinical development. Given the demands on drug discovery teams to produce molecules devoid of toxicity concerns, a paradigm shift has occurred such that toxicologists are now integral members of these teams with new responsibilities geared toward lead optimization. As such, it is no longer sufficient for the toxicologist to simply describe the observed organ toxicities of a new
compound, it is expected that we assess risk for toxicity in a predictive manner, or ascribe a mechanism to early toxicity findings, link the findings to an offending moiety within the structure, and assist in defining the chemical lead optimization path. In many instances, simple modifications to structural or physical chemical properties to molecules with insufficient safety profiles can quickly lead to new candidates with improved exposure and target organ toxicity profiles. Examples will be presented where recognition and subsequent modification of structural and physical chemical features were employed to reduce attrition of small molecules due to undesirable safety pharmacology or target-organ toxicity.

40 The Challenges of Putting In Vitro Safety Assays into Context with In Vivo Data.
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In the quest for higher-throughput, lower cost assays for safety assessment, many efforts have focused on cell-based in vitro assays that generate a effect at a certain concentration level readout such as an LC50 etc. However, most of the efforts on correlating these in vitro concentration responses to in vivo effects have looked at the simple presence or absence of a phenotypic response in the in vivo study resulting in the potential misinterpretation of a result. In this presentation, we will describe efforts to more fully describe the in vitro responses in the context of compound exposure to enhance the interpretation and correlation of in vitro assays. We will highlight some of the issues in working with in vivo data such as the loosely controlled variables used in describing the findings observed in a study and approaches we have taken to overcome these. Finally, we will show how this may then enhance the predictive value of an in vitro assay such as a simple ATP depletion assay.

41 Relating Molecular Properties and In Vitro ADME/Tox Surrogate Assay Results to In Vivo Outcomes.
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A primary goal of lead generation and lead optimization is to identify compounds with good pharmacological properties and optimal absorption, distribution, metabolism, excretion and toxicological (ADMET) characteristics. Chemists routinely consider molecular properties in designing compounds and utilize in vitro ADMET surrogate assays to select promising compounds for in vivo studies. A number of recent reports have investigated the relationship between computed molecular properties and in vivo ADMET outcomes. Although there is consensus that compound properties are important, one limitation of these analyses is a lack of controls for possible covariates (i.e., correlation vs. causation). We have examined the relationship between molecular properties and in vivo ADMET surrogate assays vs. in vitro properties within 1/5 chemical series identified from a database of 3792 compounds for which short term rodent pharmacokinetic and toxicology data are available. The role of confounding covariates was minimized by focusing on those variable associated with large differences in outcomes between compound pairs within chemically similar series of molecules. The analysis identified the following pairs of surrogates as most predictive among those examined: rat primary hepatocyte (RPH) cytolethality / volume of distribution (Vd) for in vivo toxicology outcomes, scaled microsome metabolism / calculated logP in vivo unbound clearance, and calculated logD / kinetic aqueous solubility for thermodynamic solubility. An important practical outcome of the analysis is a set of guidelines defining the utility of specific surrogates for several in vivo ADMET endpoint and the magnitude of change required in a surrogate endpoint to achieve a desired in vivo result for novel drug candidates.


Although regulated individually, the criteria air pollutants, NOx, Sox, CO, PM, Pb, and O3, exist as a complex mixture in the atmosphere. Thus, the interactions and cumulative effects of multiple pollutants are important to consider when assessing the impact of ambient air exposures on health. The criteria air pollutants act through complex biological pathways to elicit health effects, but many share common modes of action, including oxidative stress and inflammation. Mode of action for a given toxic agent is defined as the set of key events involved in a given toxic effect. Key events are measurable endpoints along a continuum from exposure to effect, and are consistent with emerging concepts of how to use biomarkers, surrogate endpoints, and toxicity pathways to characterize health risks. Recent discussions have proposed the use of mode of action to develop a unifying framework for the evaluation of mixtures containing multiple pollutants. This session convenes experts in the modes of action of criteria air pollutants to examine how this information can be organized, beginning with a description of an emerging framework for integrating mechanistic and biological plausibility information regarding the criteria air pollutants and a subset of ambient air toxics. Attendees will gain knowledge of how a mode-of-action framework can be used to consider mixtures, as well as emerging and established mechanisms of toxicity of air pollutants, focusing on potential interactions between pollutants encountered in a multipollutant context.

43 Assessing Health Effects of Air Pollution Mixtures: Mode-of-Action Framework.

To better understand the health effects resulting from exposures to mixtures of criteria pollutants (PM, particulate matter; sulfur oxides, SOx; nitrogen oxides, NOx; ozone, O3; lead, Pb and carbon monoxide, CO) that actually occur in ambient air, we are developing a framework for integrating information provided by toxicological, epidemiologic, and controlled human exposure studies based on mode of action. Mode of action for a given agent is defined as the set of key events which result in a toxic effect. Key events are measurable endpoints along the continuum from exposure to effect, and are consistent with emerging concepts of how to use biomarkers, surrogate endpoints, and toxicity pathways to characterize health risks. The approach elucidates commonalities in key events and pathways triggered by exposure to multiple pollutants. Available literature regarding the respiratory health effects of PM, SOx, NOx and O3 indicates that all four pollutants activate neural reflexes, increase bronchial reactivity, initiate inflammation and modulate immune responses which may lead to the exacerbation of allergic responses, asthma and altered host defenses. Emerging evidence also demonstrates that exposure to either PM or O3 may result in systemic inflammation, oxidative stress and impaired vasomotor function while exposure to NOx may result in pro-inflammatory circulating factors; all of which may lead to cardiovascular health effects. Exposure to CO may impair vasomotor function or lead to myocardial ischemia, while exposure to Pb may result in hypotension. Incorporation of important considerations for extrapolation in risk assessment such as adaptive responses, susceptibility factors, dose and duration of exposure, toxicokinetics, toxicodynamics and endogenous species in this framework is expected.

44 The Effects of Criteria Pollutants in the Brain.
M. L. Block. Anatomy and Neurobiology, Medical College of Virginia, Richmond, VA.

Increasing evidence links air pollution to central nervous system (CNS) disease, but the mechanisms remain poorly understood. Experimental studies and human case reports reveal that neuroinflammation and oxidative stress have emerged as a common deleterious pathway triggered by exposures to diverse criteria pollutants, including ozone and particulate matter. Thus, evidence indicates that not only does the brain’s innate immune system respond to criteria pollutants, but that this has been linked to neurodegenerative-disease-like pathology in cell culture, rodent models, and humans. This presentation will focus on the cellular and peripheral mechanisms that may be driving the CNS effects of air pollution, as our work with diesel exhaust shows that both the peripheral immune system and microglia, the brain’s resident innate immune cell, may be key to this process. In addition to providing new information on specific mechanisms of CNS disease development related to criteria air pollutant exposure, a mode of action framework will be used to describe the pathways leading to emerging health effects with uncertain mechanisms.

45 Air Pollution and Pattern Recognition Receptors: Like a Wee LCMS in Every Mouse.
M. J. Campen. College of Pharmacy, University of New Mexico, Albuquerque, NC.

Distinguishing among the toxicities of the numerous criteria pollutants and other unregulated air hazards that exist in the multipollutant environment challenges the most sophisticated human and animal studies. To adopt a more refined approach to assessing the effects of multipollutant exposures, it will be essential to better understand the shared mechanisms driving toxicity. One common pathway that several labs have recently elucidated relates to a class of receptors known as pattern recognition receptors (PRR). Research will be presented looking at the role of PRR in
Multiple epidemiological and toxicological studies have reported differential responses to fine particulate matter components, such as elemental carbon, organic carbon, sulfate, nitrate, and individual elements. While different PM components likely invoke unique pathophysiological pathways, it is also likely that there is overlap between mechanisms of action. For example, oxidative stress is a likely pathway for both organic PM components as well as trace elements. In a multipollutant setting, it is critical to understand the underlying mechanisms of adverse biological responses to these materials. This presentation will review what is currently known about pathophysiological pathways of response to PM components, with an emphasis on shared pathways.

Most seafood samples that were tested after the oil spill had dissipated, and before waters were reopened for fishing, did not contain measurable levels of oil or dispersant residues. The samples that did contain measurable residues were consistently 100 to 1000-fold below levels of concern established in the unified seafood safety protocol.

Potential toxic chemistry from the 2010 BP oil spill and clean-up efforts generated widespread public concern over the safety of seafood harvested from the Gulf of Mexico. Government agencies moved swiftly to contain the threat and reassure consumers. The effort, though, masked a larger truth. While contamination events caused by human activities draw public attention, naturally occurring toxins and deadly bacteria in Gulf waters sicken, maim and kill far more people than oil and chemical dispersants. Rare cases of neurotoxic shellfish poisoning occur in Texas. Much more common are illnesses and deaths from naturally occurring vibrios. Oysters harvested from Texas coastal waters have killed 53 people since 1995 and resulted in 45 serious illnesses from Vibrio vulnificus contamination. The presentation will cover the risks to human health from and naturally occurring contaminants, focusing on shellfish. It will also cover responses, adequate and inadequate, by governmental food safety agencies and the fishing industry.
52 Developmental Toxicity and Neurobehavioral Effects of Dietary Flavonoids.

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Flavonoids are a structurally diverse group of phytochemicals that are known hormone mimics (phytoestrogens), and are thought to have therapeutic properties. Humans are exposed to flavonoids through consumption of fruits, vegetables, and dietary supplements. These ubiquitous chemicals are found in baby formulas and foods, and are detected in human urine, plasma, and breast milk. Therefore, the potential for developmental effects should be explored. Using zebrafish (Danio rerio) as a model vertebrate, we tested the hypothesis that flavonoids exhibit structure-dependent effects on development. Embryos were treated with 5 representative flavonoids (apigenin, biochanin A, S-equol, galangin and kaempferol), 1-50 μM, from 6 hours post fertilization (hpf) to 120 hpf. At 120 hpf, effects included yolk-sac and pericardial edemas, axis curvature, fin dysmorphogenesis and craniofacial abnormalities. For all 5 compounds, we also observed spastic pectoral fin and caudal tail movements at 72 hpf, suggestive of neurotoxicity. To test the hypothesis that flavonoids are stimulants, we assessed acute effects on neurobehavior in naïve 120 hpf larvae challenged with 16 flavonoids: apigenin, biochanin A, chrysin, daidzein, dalenin, genistein, genipin, kaempferol, luteolin, myricetin, naringenin, puerarin, quercetin and resveratrol. With the exception of puerarin, all induced hyperactive swimming behavior suggesting that flavonoids have psychoactive and stimulant properties. This zebrafish larval bioassay is amenable to rapid throughput screening for anxiogenic and anxiolytic properties, which we validate using neurotoxictants chosen to represent diverse mechanisms (e.g. nicotine, chlorpyrifos, picrotoxin, etc.). Using pharmacological intervention with receptor specific chemicals, this screen will be valuable for identifying interaction of dietary flavonoids and other chemical of interest with neuro-receptors to identify the mechanism of toxicity in vivo. This research is supported by NIEHS grants F30 ES002180, RC4ES019764 and T32 ES07060.

53 Perinatal Toxicity and Carcinogenicity Studies of Styrene Acrylonitrile Trimer, a Ground Water Contaminant.

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Styrene Acrylonitrile (SAN) Trimer is a by-product of the production of acrylonitrile styrene plastics. Following the report of a childhood cancer cluster in the Tom's River section of Dover Township, New Jersey, SAN Trimer was identified as one of the groundwater contaminants at Reich Farm Superfund site in the township. The contaminants from the Reich Farm site's ground water plume impacted two wells at the Parkway well field. The National Toxicology Program (NTP) studied the toxicity and carcinogenicity of SAN Trimer in F344/N rats exposed during their perinatal developmental period and adulthood. The chronic toxicity and carcinogenicity studies in rats were preceded by 7 and 18-week perinatal toxicity studies to determine exposure concentrations for the 2 year studies. Pregnant dams were exposed to SAN-Trimer in the diet at 400, 800, or 1600 ppm during gestation, nursing and weaning periods of offspring followed by 2 years of adult exposure to both male and female pups. There was no statistically significant evidence of carcinogenic activity following SAN-Trimer exposure; however, rare neoplasms in the brain and spinal cord were noted in males and to lesser extent in female rats. These incidences were considered within the range of historical control background in the animal model used in the current studies. The major finding was a dose-related peripheral neuropathy associated with the sciatic nerves in females and spinal nerve roots in males and females, thereby suggesting that SAN-Trimer is a potential nervous system toxicant. Other non-neoplastic lesions included increased incidences of some lesions in the bone marrow and liver in males and females, and in the urinary bladder in females.

54 Does Developmental Hypothyroidism Produce Lasting Effects on Adult Neurogenesis?


The subgranular zone of the dentate gyrus (DG) of the adult hippocampus generates new neurons throughout life. Thyroid hormones (TH) are essential for brain development, but impaired neurogenesis with adult hypothyroidism has also been reported. We investigated the role of milder degrees of TH disruption on adult neurogenesis following hypothyroidism induced during development, in adulthood, or both. Pregnant dams were administered the TH synthesis inhibitor, propylthiouracil (PTU) (10 or 30 ppm in drinking water) from gestational day 6 and pups were weaned to control water on postnatal day (PN)21. On PN60, offspring from control or PTU dams were either re-exposed to PTU (30ppm) for 1 month or maintained on control, Bromoethenouridine (BrdU 50 mg/kg, ip, twice daily) was administered to all animals on the last 5 days of the re-exposure period, and animals sacrificed 28 days later. Animals were perfused transcardially, the brains removed and embedded in a MultiBrain (NSA) array and freeze sectioned. Every 8th section throughout the hippocampus stained with an antibody against BrdU to mark actively dividing cells. The volume of the DG and the number of BrdU-positive cells were assessed from images captured on a Nikon microscope (400X) and Nikon Elements software. Preliminary findings indicate that developmental exposure to PTU produced a persistent reduction in the volume of the adult DG. BrdU cell counts were reduced similarly in all PTU-exposed groups. These data suggest that moderate levels of hypothyroidism decrease cell survival in the adult brain and that transient developmental hypothyroidism leads to persistent decreases in DG volume and cell survival. The degree to which these findings are determined by reductions in cell proliferation is currently under investigation. As neurogenesis in the adult recapitulates developmental processes of proliferation, differentiation, and migration, study of this neurogenic niche in the adult may provide a simpler means to assess the consequences of TH insufficiency on neurodevelopment. (Does not reflect EPA policy).


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A number of publications have reported that i.p. administration of paraquat (PQ) to rodents (e.g. C57Bl6) mouse) at high doses results in a loss of dopaminergic neurons from the substantia nigra pars compacta (SNpc), the primary area of neuropsychological damage in Parkinson's disease (PD). Such studies have been used to indicate mechanistic plausibility for epidemiological claims of a link between PQ exposure and PD. A major criticism of the i.p. mouse model is that it uses a route of administration not expected with PQ which is absorbed into the skin. To better understand the relevance of the reported findings from the i.p. mouse model we have conducted a study where male C57Bl6 mice were exposed to PQ in the diet for 13 weeks, and the brains examined for evidence of dopaminergic neuronal cell loss using stereology, changes in striatal neurotransmission and pathological changes using stains to detect neuronal cell damage and inflammatory responses. Dietary concentrations of 10 & 50 ppm paraquat dichloride salt were used which resulted in achieved doses of 2.4 & 14.1 mg/kg/day. A low dose of N-1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP; 10 mg/kg administered i.p. to a separate group of male mice 4 times in a single day at 2 hr intervals, 7 days prior to the end of the dietary study) served as a positive control. PQ at either dose level did not induce a loss of tyrosine hydroxylase positive (TH+) dopaminergic neurons in the SNpc, after the concentration of striatal dopamine and its metabolites or result in evidence of neuronal cell damage or astrocyte/microglial activation in the SNpc. In the MPTP group the number of TH+ neurons in the SNpc and striatum decreased; however there was significant pathological changes including neuronal necrosis and astrocyte/microglial activation. This study further brings into question the relevance of the findings from previous i.p. mouse studies as evidence for a link between PQ exposure and PD/parkinsonism.

56 Neurodevelopmental Effects of Inhaled Vapors of Gasoline and Ethanol in Rats.

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Gasoline-ethanol blends comprise the major fraction of the fuel used in the US automotive fleet. To address uncertainties regarding the health risks associated with exposure to gasoline with more than 10% ethanol, we are assessing the effects of prenatal exposure to inhaled vapors of gasoline-ethanol blends. Pregnant Long-Evans rats are exposed to fuel vapors, 6.5 hr/day, on days 9–20 of gestation, and their offspring are assessed for a variety of neurodevelopmental effects. This report compares effects of inhaled gasoline vapor lacking ethanol (E0) at concentrations of 0, 5000, 10000 or 21000 ppm with previously-reported effects of inhaled ethanol (E100) at concentrations of 0, 5000, 10000 or 21000 ppm. Maximum concentrations were limited by the lower explosive limits of the vapors. As observed with
E100, E0 vapors caused no overt maternal toxicity, changes in litter size or weight, or weight gain of the pups. In contrast to E100, E0 did not alter locomotion activity of adults. Both E0 and E100 produced a few minor, unsystematic changes in the functional observational battery. In water maze tests, E100 altered search strategies (not dose-related) and impaired memory in females (no-platform probe trials, all doses); in contrast, E0 was essentially ineffective. No treatment affected working memory, as assessed by the object delayed-matching-to-position tests. In an operant reaction-time test, 21000 ppm E100 and 9000 ppm E0 increased hold failures, a measure of impulsivity, and increased decision times at the lowest doses. Responses to a conditioned audiovisual cue were reduced in females at all doses of E100, but were not affected by E0; neither agent affected conditioning to context. Telemetered mean blood pressure (BP) was increased in male offspring by 9000 ppm E0 at PND 90 and 180; tail cuff tests corroborated these results at PND 180 only. E100 increased BP (tail cuff) in males at all doses on PND90 only. This abstract does not reflect EPA policy.

**57**

Postnatal Trichloroethylene Exposure Is Associated with Abnormal Behavior and Alterations Global DNA Methylation Patterns in Mouse Cerebellum.


Previous studies have shown that continuous exposure throughout gestation until the juvenile period to environmentally-relevant doses of trichloroethylene (TCE) in the drinking water of MRL+/- mice promoted adverse behavior associated with glutathione (GSH) depletion in the cerebellum indicating increased sensitivity to oxidative stress. Here we extend these findings to further characterize the impact of TCE exposure on redox homeostasis, biomarkers of oxidative stress, transmethylation metabolites and global DNA methylation patterns in mice exposed to water only and two doses of TCE in the drinking water postnatally from birth until 6 weeks of age. The mice were subjected to a variety of open field behavioral tests in order to correlate behaviors with metabolic and methylation patterns observed in our model. Our results show that the cerebellum from male mice exposed to TCE have lower GSH and increased markers of oxidative stress compared with controls. Methionine levels were also significantly reduced in the TCE-exposed mice which suggested compromised cellular methylation. Global DNA methylation, including hydroxymethylation patterns, were significantly lower in the cerebellum of TCE-exposed mice, compared to controls. Mice exposed to TCE exhibited increased locomotor and exploratory activity compared to control mice suggesting increased anxiety/safety seeking behavior similar to that observed in humans with attention deficit disorder. Understanding the mechanisms of TCE neurotoxicity during sensitive windows of exposure is important in order to enhance mechanistic understanding of environmentally-related neurologic disorders in susceptible populations.

**58**

Perfluorohexane Sulfonate (PFHxS) Causes Adult Behavioral Disturbances in Male and Female Mice, after Neonatal Exposure.


Perfluorohexane sulfonate (PFHxS) is a perfluorinated compound (PFC) used as an industrial additive. PFC chemical properties make them suitable as surfactants and oil- and water repellents, which are frequently used in products for packaging and as protective coatings. However, the same properties also account for their extreme physico-chemical stability, making them practically non-biodegradable, accumulat- ing in the global environment, causing concern, since little is known about the toxicity of the compound. We recently have seen that other PFCs, like perfluorooctanoate sulfonate (PFOS) and perfluorooctanoic acid (PFOA), can induce developmental neurotoxic effects and the purpose of the present study was to explore if neonatal exposure to PFHxS can affect behavior and cognitive function. In the present study we exposed male and female mouse pups to a single dose of PFHxS (0.61-9.2 mg/kg bw) during the defined critical period of brain development, on postnatal day 10. At two months of age male and female mice showed altered spontaneous behavior in a novel home environment, affecting cognitive function. Furthermore, these functional behavioral effects were long-lasting or irreversible since they were once again seen at four month of age. The nicotine-induced behavior test revealed that male and female mice neonatally exposed to PFHxS responded differently compared to control animals, when challenged with a dose of nicotine. The present findings show that PFHxS can cause developmental neurotoxicity effects similar with effects earlier reported after neonatal exposure to PFOS and PFOA and other persistent pollutants, such as PBDEs and PCBs.

**59**

Acute Ozone-Induced Impairment of Glucose Regulation: Age-Related and Temporal Changes.


Ozone (O3) is associated with adverse cardiopulmonary effects in humans and thought to produce metabolic effects, such as insulin resistance. We showed that episodic O3 exposure increased insulin levels in aged rats. We hypothesized that O3 could impair glucose homeostasis by altering insulin signaling and/or causing an unfolded protein response (UPR) in the liver. Brown Norway rats, 1, 4, 12, 21, and 24mo old, (a model of non-obese aging) were exposed to O3 at 0.5, 0.5ppm, and 6h/day for 2 consecutive days. As a follow-up study, 4mo old rats were exposed to 0 or 1ppm O3 over 2 days to examine the time course of response. Glucose tolerance tests (GTT) directly followed exposure in all studies and additionally at 24h post-exposure in the time-course experiment. Liver gene expression was examined using Affymetrix RG-230PM Array strips. Liver and adipose tissues were also analyzed using RT-PCR for metabolic and UPR markers. Phospho-protein analysis to assess insulin signaling was done in the liver, adipose tissue, and muscle. GTT showed a marked impairment of glucose clearance among 1ppm O3 exposed rats of all ages. The reduction in glucose regulation after O3 exposure was most apparent in 1mo rats, who exhibited no baseline glucose intolerance. The 24mo rats exhibited glucose intolerance at baseline. Analyses of metabolic and acute phase response (APR) biomarkers show that the insulin signaling pathway is altered by O3 in all three tissues and serum APR proteins were increased. Selected UPR genes were upregulated in the liver. Serum leptin increased acutely following 1day (6h) O3 exposure. Maximum effects of O3 on insulin signaling and APR were seen directly following the second day of exposure. Our results suggest that glucose intolerance is a result of metabolic changes in response to O3. These findings raise concern about ambient O3’s potential to cause predisposition towards metabolic impairment. (Does not reflect US EPA policy).

**60**

Fine Particulate Matter (PM2.5) Exposure Impairs Vascular Insulin Signaling and Exacerbates Diet-Induced Systemic Insulin Resistance in Mice.

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Recent epidemiological studies suggest that increases in fine particulate matter (PM2.5) air pollution contribute to the rapidly evolving epidemics of obesity and diabetes. Because metabolic syndrome, diabetes and air pollution all induce endothelial dysfunction, and because we recently showed that short-term exposure to centimeter ambient PM2.5 (CAP) improves endothelial VEGF signaling and decreases circulating endothelial progenitor cells (EPCs), we examined the effects of CAP exposure (9-30 consecutive days, 6h/d) on endothelial and systemic insulin resistance and inflammation, adiposity and vascular function in mice fed normal chow or high-fat diet (HFD). Surprisingly, CAP exposure impaired vascular insulin signaling after only 9 or 30 days, and exacerbated HFD-induced systemic insulin resistance after 30 days without increasing adiposity. Insulin sensitivity and inflammation in adipose and liver were unaltered by CAP or HFD. In contrast, CAP exposure (30 days) impaired endothelial insulin signaling diet-independent in heart and aorta, but HFD-dependent skeletal muscle. Changes in insulin sensitivity were accompanied by organ-specific but not systemic inflammation and a decrease in circulating EPCs. Collectively, our results suggest that short-term CAP exposure provokes HFD-induced systemic insulin resistance by inducing inflammation and endothelial insulin resistance accompanied by decreased circulating EPC levels. Impaired vascular maintenance due to EPC suppression could contribute to vascular insulin resistance (or vice versa) thereby increasing systemic insulin resistance and the risk for the development of T2D and CVD by PM2.5.

**61**

Ozone (O3): A Potential Contributor to Metabolic Syndrome through Altered Insulin Signaling.


Air pollutants have been associated with diabetes and metabolic syndrome, but the mechanisms remain to be elucidated. We hypothesized that acute O3 exposure will produce metabolic impairments through endoplasmic reticular stress (ER) stress

**SOT 2013 ANNUAL MEETING 11**
Inhaled Ozone Induces Metabolic Abnormalities in Mice Fed a High-Fructose Diet.

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Results of recent inhalation toxicology studies have indicated that long-term exposure of mice to concentrated ambient particulate matter leads to increased insulin resistance and potentiates other adverse metabolic effects brought on by consumption of a high fat diet. Similar adverse metabolic effects on insulin signaling and glucose homeostasis caused by gaseous air pollutants, such as ozone (O3), have not been investigated. In the present study, we tested the hypothesis that subacute inhalation exposure to O3 enhances metabolic abnormalities, caused by a high fructose diet (HFrD), in mice. C57/Bl6 male mice were maintained on either a normal chow diet (NC) or a HFrD (from which 60% of the calories were derived from fructose) for 8 weeks prior to exposure and during exposure. The purpose of this study was to characterize the cardiovascular effects of O3 exposure in mice fed a high fructose diet (HFrD). Rats were exposed once for 3 hours to 3 parts per million acrolein gas or filtered air (control) and arterial blood samples were obtained. Arterial blood samples were obtained immediately following each day or one day after 2-day exposure. Tissues were analyzed for insulin and ER stress signaling and serum for inflammation and metabolic biomarkers. O3 produced severe hyperglycemia and glucose intolerance that was reversible following 1 day recovery. Phosphorylation of insulin receptor substrate (pIRS) decreased after 2nd day O3 in all three organs. Downstream mediators, phospho-serine/threonine kinase and phospho-glycogen synthase kinase also decreased but only in adipose tissue. Serum insulin changes were correlated with tissue levels of pIRS. Serum IL-6, thought to link ozone-induced inflammation and metabolic alterations, did not increase at any time; however lipocalin and acute phase proteins increased after O3. Serum leptin was also increased sharply after 1-day O3 exposure and it was correlated with O3-induced hyperglycemia, but not glucose intolerance. Genes downstream of unfolded protein response were changed in the liver indicating ER stress. To examine the role of liver ER stress in O3 induced impairment of metabolism, we treated rats with ER-stress inhibitor, salubrinal prior to air or 1ppm O3. Salubrinal did not diminish O3-induced hyperglycemia and glucose intolerance, suggesting that hyperglycemia likely did not result from ER stress. In conclusion, acute O3 exposure alters insulin signaling in metabolic organs causing hyperglycemia and glucose intolerance, which might contribute to increased incidences of diabetes and metabolic syndrome.

Pl 63 Suppressed Responses in Heart Rate Variability during Inhalation Exposure to Ozone and Ambient Fine Particles in Rats on a High-Fructose Diet.

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People with diet-induced cardiometabolic disorders (diabetes, hypertension) may be more susceptible to the adverse cardiovascular effects of air pollution. The present study was designed to determine if a high-fructose diet affects cardiovascular responses to inhalant air pollutants. We used a mobile air research laboratory located in an industrial area of Dearborn, MI, to expose male Sprague Dawley rats, to filtered air (FA), ozone (O3; 0.5 ppm), concentrated ambient fine particles (CAPs; 400 μg/m3) or the combination of O3 & CAPs. Rats were fed a normal (ND) or high-fructose diet (HFD) for 8 weeks prior to exposure and during exposure. Inhalation exposures were 6h/day for 9 days (4-5 days/week). Heart rate (HR) and ECG waveforms were collected by radiotelemetry every 5 minutes during exposures, and measures of heart rate variability (HRV, indicated as SDNN and rMSSD) were calculated and analyzed by a linear mixed model. Compared to FA-exposed HFD, HFD had significantly greater HR (328 ± 27.5 vs 299 ± 35 bpm, respectively) and lower SDNN (44.6 ± 78.8 vs 64.4 ± 86 ms). All exposures caused decreases in HR that were greater in HFD rats (25-40 bpm) than in ND rats (6-15 bpm). However responses in HRV were dependent on the specific exposure and diet. Exposure to O3&CAPs induced significant decreases in HRV in ND- but not HFD-fed rats, with a 55% decrease in SDNN over the nine days of exposure. By comparison exposure to O3 alone had the opposite effect on HRV, with a 48% increase in SDNN, but with no effect in HFD rats. Lastly, ND-fed rats exposed to CAPs alone had a modest decrease in SDNN (23%), while similarly exposed HFD-fed rats again had no change. In summary, while HR was decreased in all experimental groups, the degree and direction in change of HRV differs on the specific exposure and diet. The mechanism(s) underlying altered autonomic responses in HFD rats to single- or multi-pollutant exposures requires further study.

Pl 64 Oxidative Stress and the Acceleration of Atherosclerosis in Susceptible Mice after Exposure to Semivolatile Components of Ultrafine Particulate Matter.

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Exposure to ultrafine particulate matter (UF-PM) has been associated with adverse cardiovascular health effects. UF-PM contains semi-organic volatile substances (SVOCs) that are bound to particles but can partition to the vapor phase after emission. SVOCs contain species such as polyyclic aromatic hydrocarbons and quinones that can induce oxidative stress and may contribute to the oxidative modification of atherosclerotic disease by UF-PM. Therefore, we hypothesized that the removal of SVOCs from an aerosol should decrease the ability of the particle to cause oxidative damage and consequently the acceleration of atherosclerotic plaque formation. ApoE -/- mice, which are prone to developing atherosclerosis, were exposed to UF-PM containing ambient particles (CAPs), CAPs with the SVOC components removed, or SVOC components without the particle core. A control group was exposed to purified, filtered air. Particles were concentrated using a VACES, and SVOCs were separated from the particle core using a thermal denuder. The exposures took place 5 hours/day, 4 days/week for 8 weeks in downtown Los Angeles, 100m downwind of a major freeway. Plaque formation in the aortic arch and total and LDL cholesterol in the serum were measured to evaluate the progression of atherosclerosis. Serum concentrations of lipid peroxidation, protein carbonyl content, and glutathione were assessed to determine systemic oxidative stress. Aortic plaque formation in mice exposed to unmodified CAPs was higher than in those exposed to CAPs with no SVOCs. Similarly, higher levels of lipid peroxidation were measured in mice exposed to unmodified CAPs and SVOC components of CAPs compared to those exposed to CAPs without SVOCs. The corresponding trends in plaque formation and lipid peroxidation suggest the tendency that exposure to SVOCs may contribute to the acceleration of atherosclerosis via an oxidative stress pathway.

Pl 65 Acrolein-Induced Increases in Blood Pressure and Heart Rate Are Coupled with Decreased Blood Oxygen Levels during Exposure in Hypertensive Rats.

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Exposure to air pollution increases the risk of cardiovascular morbidity and mortality, especially in individuals with pre-existing cardiovascular disease. Recent studies link exposure to air pollution with reduced blood oxygen saturation suggesting that hypoxia is a potential mechanism that mediates the adverse cardiovascular effects of air pollution. The purpose of this study was to characterize the cardiovascular effects of exposure to acrolein, a potent irritant and component of cigarette smoke and diesel exhaust and determine if acrolein exposure causes decreased blood oxygen levels. We hypothesized that hypertensive rats would be more sensitive to the adverse cardiovascular effects of acrolein and that the cardiovascular effects of acrolein would be coupled with decreased arterial blood oxygen levels during exposure. Spontaneously hypertensive (SH) and Wistar Kyoto (WKY; rats with normal blood pressure) rats implanted with biopotential radiotelemetry transmitters were exposed once for 3 hours to 3 parts per million acrolein gas or filtered air (control) in whole body plethysmograph chambers while cardiovascular and ventilatory parameters were monitored. In a separate cohort of rats, arterial blood samples were drawn before, during, and after exposure to acrolein to monitor blood oxygen saturation. We found that hypertensive, but not normal rats, had significant increases in
Diabetes are particularly vulnerable to the adverse cardiovolumetric effects of particulate air pollution (PM)—often attributed to enhanced inflammation and endothelial dysfunction. Recent reports have implicated activated receptors for advanced glycation end-products (RAGE) as an integral factor in the inflammatory processes of cardiovascular dysfunction and diabetes; nonetheless, it is unclear whether ambient PM alone/or in combination with other endogenous factors may contribute to RAGE activation. Levels of soluble RAGE (sRAGE) were measured in human serum samples (n=60) from two cities in Gandhi, China (Jingcheng (JC) and Zhanpeng (ZH)—albeit similar ambient levels of PM; NI, Cu, As, and Se in JC were 76, 25, 17, and 7 fold higher than ZH, respectively). In addition, human pulmonary endothelial cells were exposed to PM (collected from the same region) to investigate RAGE-mediated vascular dysfunction. Lastly, to examine the link between PM and overt diabetic endpoints, B6C3/F2 mice were exposed to concentrated ambient PM (CAPS). Results: sRAGE was significantly higher (p<0.04) in residents of JC (538.7±37.5ng/ml) than that measured in ZH (452.6±21.6ng/ml). Multiple regression analyses revealed PM2.5 concentration as a significant (p<0.03) predictor of RAGE outcome. In the in vitro work, after 48h of PM2.5 exposure, a dose dependent increase in cell proliferation and small increases in sRAGE activity at higher doses of PM was evident. Immunofluorescence detection showed an elevation in cells positive for membranous RAGE expression; accompanied w/ a 2-fold increase in mRNA for RAGE & NF-kB and >2-fold increase of ATF4 & NF-kB in cells treated w/ PM-BSA. These findings suggest plausible interaction between PM & RAGE resulting in enhanced expression of NF-kB, ATF4 and RAGE. Finally, preliminary mouse exposures have yielded supportive findings: progeny of CAPS exposed pregnant mice have shown significantly decreased glucose tolerance compared to controls (p<0.03). Collectively, these data offer valuable insight into PM-mediated RAGE activation / its influence on diabetes.

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Integration of telemetered hemodynamics in toxicology studies is an emerging trend which offers detailed information on potential cardiovascular safety issues that are only identified with repeat dosing. Jacketed external telemetry (JET) is an enabling technology for which the intrinsic variability of the associated minimally invasive blood pressure (BP) device (PA-C10, DSI, St. Paul, Minn.) has not been extensively characterized. Accordingly, we evaluated the intrinsic and extrinsic beat-to-beat measurement error of JET BP in 10 male cynomolgus monkeys (2.7-3.7 yrs, 3.0±0.2 kg). BP catheters were advanced to the descending aorta via the femoral artery. BP (24 h) was continuously digitized (500 Hz) in a quiet radio frequency environment and retrospectively analyzed in beat-to-beat increments (Life Sciences Suite, ver. 5.0, DSI). Data are presented as mean±std. Sham dosing occurred at 2.5 h post acquisition start to mimic a typical toxicology study design. Errorous values were segregated as beats where either systolic pressure, pulse height, and/or <p>Dp/dt were 60c>240, 20c>80, or 600c>4000 native units, respectively. Transducer stability was assessed as the beat-to-beat change (Δ) in systolic pressure. Over 24 h, systolic pressure was 130±8.5 mmHg and Δ was 0.02±2.67 mmHg. During a quiescent nocturnal period (10 h) systolic pressure was 131.9±8.5 mmHg and Δ was 0.02±2.42 mmHg. Errorous values (8574/4246±2 beats, 0.36%) were infrequent and temporally random. These data confirm that the JET BP device conformed to, or exceeded, design sensitivity (±3 mmHg) at all times. Systolic pressures exhibited stable diurnal variations of ±8.5 mmHg yielding > 80% power to detect a 10 mmHg change (p<0.05) with n=6. The sparse, sporadic incidence of non-physiologic hemodynamic values was insufficient to influence mean BP values aggregated over times ≤ 5 min. Consequently, resources devoted to the beat-to-beat filtering of JET BP values will not improve the accuracy, precision, or power of these data.

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EMG noise and movement artifact are barriers to obtaining accurate interval measurements in ambulatory subjects in preclinical studies. Herein we report the results of testing a new device, an In Line Filter (IF) that employs Multi-Domain Signal Processing™ (MDSP) technology, for real-time noise and artifact suppression in telemetered subcutaneous ECG recordings. Prior evaluations of MDSP technology using ECGs already acquired by the Dataquest™ system showed that it reduces the amplitude of in-band noise by > 95% without distorting ECG morphology and provides a corresponding increase in the number of analyzable beats. In this work, studies of ILF performance were conducted using freely moving subiects instrumented with DSI™ D70-PCT telemetry transmitters. An ILF was inserted between the telemetry receiver and the Dataquest system to filter out noise. Raw and filtered ECG signals were collected from multiple species commonly used in safety studies and were analyzed with emka ECGauto using standard methods. Results obtained from filtered and unfiltered recordings were evaluated to assess the impact of filtering on parameter measurement accuracy, the increase in beats available for analysis, and the amount of labor required to perform interval analysis. Results show that, on average in all species tested, filtering the ECGs using the ILF prior to analysis increased the analyzable portion of the data to roughly that reported in the literature for telemetered epicardial and intravenous ECG sensors and significantly reduced the amount of labor required to perform interval analysis. There was no degradation in the accuracy of parameter measurements as a result of filtering the signal. Use of the ILF could make routine beat-to-beat analysis of 24-hour subcutaneous ECG recordings practical and cost-effective without the complications and complexities associated with epicardial, endocardial, and intravenous ECG sensing leads.


The species differences are the common and major concerns in safety assessment and translational science, particular in situations where results from CV and CNS safety studies in the same species are required or the receptor subunits expression are significantly difference cross species etc. In this study, we have used the radio-telemetry technique to record ECG and EMG form conscious, non-sedated dogs and monkeys and validated the model with a proconvulsant, pentylenetetrazole (PTZ). Beagle dogs and cynomolgus monkeys were implanted with four electrodes epideral over the motor cortex and connected to the radio-transmitter (DSI) placed subcutaneously in the lateral neck region. Two additional electrodes were inserted into the ridge cervical muscle for EMG recording simultaneously. The signals captured by the receiver were routed via a data matrix board to a PC and sampled at a rate of 500Hz. Subcutaneous (s.c) administration of PTZ dose-dependently (10, 20 and 50 mg/kg s.c) induced paroxysmal activity, clonic convolution and tonic convolution in both dogs and monkeys, indicating a remarkable CNS safety issue of PTZ as positive control article. When PTZ administrated i.v at 1.5mg/kg/min and 1.5mL/min, both dogs and monkey showed similar level of paroxysmal spike-and-wave activity associated with clonic convulsions occurred between 17 and 36 min after the start of infusion at 4.5Hzs. The data indicate that the electroencephalogram (EEG) and electromyogram (EMG) are the most sensitive and valuable biomarkers in identifying pathologic CNS activity, in particular for safety pharmacology evaluation and translational application.

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A critical aspect of cardiorespiratory safety pharmacology studies is the collection of high quality cardiovascular and respiratory data immediately prior to dose, during dose and immediately after dose to be able to recognize any acute effects of the test
article. This is especially challenging when the test article is administered via inhalation. This poster describes the method for achieving a successful outcome in these types of studies, including the factors that must be considered in the design and execution. Effects of proper habituation to equipment and carefully scheduled study activities were assessed based on the overall character of cardiorespiratory response from 3 different studies. Prior to the first data collection, all animals were surgically implanted with DSI telemetry transmitters and habituated to the exposure/data collection system. Respiratory parameters were collected using respiratory inductance plethysmography (RIP). Telemetry data were collected continuously while animals were in their home cage, after transfer to the exposure suite but prior to either air or the vehicle, during the exposure period, and for 24 hours post dose in 3 different studies. Each study used a different vehicle and an air ( sham) control was administered to assess the vehicle effect. Administration of the vehicle produced no effects on blood pressure, heart rate, body temperature, respiratory rate, tidal volume and minute volume during pre-exposure period and for the 24 hour post dose recording period when compared to air ( sham) control in all 3 studies. Very similar patterns in blood pressure and heart rate were noted in all 3 studies; however, each study has a specific pattern for body temperature and respiratory parameters, which was attributed to individual variation in these parameters. This poster presents a consistent and reliable method of collecting cardiorespiratory data from conscious beagle dogs prior to, during, and after inhalation exposure in our Testing Facility.

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The detection of delayed ventricular repolarization, characterized by a QT interval prolongation, is one of the main issues for the preclinical evaluation of potential risk of pro-arythmia for a new drug candidate. This study was designed to compare different methods of QT interval prolongation assessment in conscious beagle dogs using external telemetric device. The same 6 dogs (3/sex) were given vehicle or reference compounds known to induce QT interval prolongation using a sequential design (single oral dosing of Sotalol at 30 mg/kg and Moxifloxacin at 30 then 90 mg/kg or repeated oral dosing for 6 days of Thoridizidine at 20 mg/kg/d and Terfenadine at 30 mg/kg/d). Electrocardiograms were recorded during a treatment-free period and on day 1 and day 6 for approximately 20 h post dosing. QT intervals were measured from a beat to beat analysis over 10-min periods centered on selected timeslots and corrected according to the Van de Waters formula (QTvdw). QT shift calculations and an analysis based on the principles of the Holzgrefe’s probabilistic model (QTh, using a minimum 250 beats/timeslot) were also performed. All methods allowed an accurate evidence of QT effect for Sotalol and Moxifloxacin. The QT shift and QTh methods gave more evidences for Terfenadine effect than QTvdw. As expected, the marked increase in heart rate (HR) induced by Thoridizidine resulted in no apparent effect on QT and a statistically significant overcorrection for QTvdw whereas QT shift and QTh gave more accurate and reliable results. The statistical sensitivity threshold detection of QT prolongation was low, i.e. 10 to 15 ms for QT shift and QTh methods. Thus, external telemetry in non-clinical safety dog studies allows new perspectives for a better assessment of QT prolongation when associated to QT shift and Holzgrefe’s probabilistic methods, especially in case of slight drug-induced QT effect or marked changes in HR.

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Integration of Automated Patch Clamp Systems and Logistic Models in the Cardiochannelgramm (CCGMT) for Better Prediction of Cardiac Risk.

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Drug-induced inhibition of the cardiac hERG potassium channel is recommended by ICH and FDA to predict delayed cardiac repolarization (DR) and cardiac risk. The consequent QTc prolongation is a surrogate marker of Torsade de Points (TdP), a rare but potentially lethal ionicotropic outcome. Drugs with effective thera-peutic plasma concentrations (ETPC) within 30-fold of their hERG IC50s are thought to be dangerous despite the fact that multiple ion channel effects (MICE) can mitigate DR. Here we demonstrate that logistic regression models, which integrate MICE, predict TdP with much greater certainty than the hERG safety ratio (hERG IC50/ETPC or safety margin (SM)) alone. To this end we measured hERG, Nav1.5,Cav1.2, Kir2.1 and Kv14QT1/minikIC50 values of 56 drugs (33 +TdP and 23 -TdP) from multiple classes using automated patch clamp systems including Qpatch and PatchXpress. The sensitivity of the automated patch clamp platforms was evaluated in a comparison to manual patch clamp. ETPC values and the torsadogenic liability of drugs was obtained from the literature, package inserts and Arizona CERT. Eleven logistic regression models were constructed; one using the hERG SM alone, the others integrating hERG SM, Nav1.5 SM and/or Cav1.2 SM data. The predictive power of each model was evaluated using the likelihood ratio test. Leave-one-out cross validations were performed and each model’s accuracy was determined by comparing receiver−operating characteristics (ROC, sensitivity vs. 1-specificity). Models that include Nav1.5, Cav.1.2 or both variables are statistically significantly better predictors of TdP liability than the model that contains only hERG (Model 1). Model 1 had a ROC area under the curve (AUC) of 0.80 and Model 11, that includes hERG, Nav and Cav.1.2 SM, significantly improved accuracy showing a ROC AUC of 0.94. Thus, Model 11 that incorporates the concept of MICE in the CardiochannelGramm (CCGMT) is a robust nonclinical predictor of cardiac risk.

P73  
Characterization of Jacketed External Telemetry with Blood Pressure in Conscious Nonhuman Primates in Pen-Style Housing Administered Etifline, Sotalol, or Hydralazine.

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An important component of nonclinical safety assessment is the evaluation of electrocardiography (ECG) and hemodynamic parameters. Jacketed external telemetry with an implanted telemetry blood pressure transmitter (JET-BP) is a minimally invasive technology being utilized in general toxicology studies to assess ECG and blood pressure measurements in conscious, unrestrained animals. We characterized JET-BP in nonhuman primates (NHPs) housed in pen-style cages with three reference compounds with known effects on blood pressure or ECG parameters. Thirty-six male NHPs were implanted with a telemetry blood pressure device and group-housed in pen-style caging (3 animals/pen). Four animals per group were given reference material or a concurrent control article. Etifline, a sympathomimetic, was given at 1 or 10 mg/kg; hydralazine, a vasodilator, at 1 or 10 mg/kg; and sotalol, a non-specific β-blocker, at 3 or 30 mg/kg. JET-BP measurements were recorded for at least 90 minutes prior to dosing and continuously for at least 20 hours postdose. The following ECG and hemodynamic parameters were determined as one (light phase) or two-hour (dark phase) averages: PR, QT and rate-corrected QT; systolic, diastolic, and mean arterial pressures; heart rate; and arterial pulse pressure. In addition, blood was collected at seven postdose timepoints for each test article for pharmacokinetic analysis. Etifline significantly increased mean arterial pressure, systolic blood pressure and pulse height, while diastolic blood pressure remained unchanged. Sotalol significantly prolonged QT with no significant change in blood pressure. Administration of hydralazine did not significantly change blood pressure at the dose levels administered. In summary, ECG and blood pressure changes caused by three different reference compounds were detectable using JET-BP technology in group-housed NHPs.

P74  
Differential Cardiovascular Physiology and Pathology in Selected Lineages of Miniature Swine.

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The miniature swine has been increasingly recognized as a valid alternative to canine and non-human primates in regulatory toxicity. This poster presents the results of cardiovascular assessments in the Yucatan, Hanford, and Sinclair miniature swine conducted during clinical investigations and control toxicity testing. Anatomic parameters were obtained at necropsy. Blood vessels diameter, velocity, and flow were obtained by Doppler ultrasonography. Cardioelectric physiology was obtained using clinical ECG and surgical monitor units. Macroscopic lesions and histopathology assessments were conducted on heart and kidneys. Data were compared to published measurements of adult human illustrating similarities or differences (for practicality, male data are reported here). Across the three lineages, heart- to-body weights ratio ranged from 0.41 to 0.50 and were higher than human (0.42). The geometric correlations for heart rate adjustments to body size ranged from 215 to 297 and were comparable to human (241), indicating that heart volume and function were well adjusted to the reduction in body size. The miniswine
hearts showed a coronary artery distribution comparable to human. The right coro-
nary internal diameters ranged from 1.44 to 1.79 mm and were comparable to
human (3.9 mm) when adjusted to body surface area (weight range: 10-30 kg).
External femoral blood flows at rest averaged 93 mL/min and were slightly lower
than human (260 mL/min) when adjusted to body size. Electrophysiological heart
segments duration (e.g. RR ranged from 360 to 662 msec) and their ratio (QT/RR)
were proportional to human and well-adjusted to body size. Macroscopic lesions
were nonexistent. Histopathology findings were rare and limited to sub-level my-
ocardial inflammation with low incidence in the Hanford lineage. In conclusion,
the similarities between the cardiovascular systems make these three lineages of
miniature swine suitable animals to model the human counterpart.

Whole Heart Electrophysiology and Stress Test As an Indicator of
Drug-Induced Cardiac Toxicity.
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Cardiac toxicity, manifested as compromised contractility or ischemic heart disease,
comprises 26.9% of post-approval drug failures. The heart has a high demand for a
constant energy supply which can be affected by many sources of stress and thus
may be a good indicator of potential toxicity. The purpose of this research was to
utilize the ex vivo heart model to assess contractility and whole heart energetics in
response to drugs with known/unknown mechanisms of toxicity. We used FCCP
and verapamil as positive and negative controls and doxorubicin (dox), doxorub-
icin-ol, sunitinib, and sorafenib as the chemotherapy agents associated with latent
toxicity. Rat hearts were removed, perfused with Modified Henseliet Krebs, and LVP
was monitored via insertion of a fluid-filled balloon. The perfused heart was in-
serted into an 11.7 T NMR magnet. Whole heart phosphorus content was assessed
before and during 60 min of drug exposure and then during 20 min of 0.1μM iso-
proterenol (iso) with drug to assess energy reserve. Control heart contractility and
energetics were stable throughout the experiment until the iso challenge, where
contractility increased as expected and PCR and ATP decreased and Pi increased.
FCCP treated hearts showed a decline in contractility and PCR and reduced reserve
during the iso challenge. Verapamil treated hearts did not change in energetics dur-
ing treatment or during the iso challenge. Dox increased contractility, while the
other chemotherapies showed very little change in contractility during drug treat-
ment. Dox treated hearts showed a drop in PCR and reduced reserve
during the iso challenge. Verapamil treated hearts did not change in energetics dur-
ing treatment or during the iso challenge. Dox increased contractility, while the
other chemotherapies showed very little change in contractility during drug treat-
ment. Dox treated hearts showed a drop in PCR and reduced reserve.

Simultaneous Recording of Action Potentials and Calcium
Transients from Stem-Cell Derived Cardiomyocytes:
Applications for Cardiototoxicity Testing.
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Beerse, Belgium; 3Sanford-Burnham Medical Research Institute, La Jolla, CA.
Current methods for preclinical cardiotoxicity testing generally examine the effects
of candidate compounds on the activity of single ion channels using manual or pla-
nar patch clamp methods. Limitations in these assays require that the tests be per-
formed with cell lines which stably express the ion channel of interest. This reduc-
tionist approach grossly underestimates the complexity of cardiomyocyte
excitability and physiology. We have developed a new automated image cytomter
and associated software which facilitates a more physiologically relevant test of
compound effects on cardiomyocyte excitation-contraction coupling. This new ap-
proach utilizes a dual channel automated Image cytometer that allows for simulta-
eous measurement of the cardiomyocyte action potential and calcium transient
using voltage and calcium sensitive dyes. By using these advanced imaging tech-
niques this system frees the need for the assay apparatus to interact physically with
the cells, allowing the use of a wide array of cell types including more clinically rele-
vant models such as cardiomyocytes derived from human induced pluripotent
stem cells (hiPSC). Here we demonstrate the application of this system to a small
scale screen of knowncardioactive compounds in hiPSC derived cardiomyocytes.
Our results suggest that the ability to identify perturbations in the cardiomyocyte
action potential and/or calcium transient due to exposure to cardioactive com-
ounds is on par with existing technologies. However, this system demonstrates a
much higher throughputs than existing systems and provides a more complete
analysis of compound effects on excitation-contraction coupling in the cardiomy-
ocyte.

Effect of Cell Culture Media on the Growth and Viability of
Neonatal Rat Cardiomyocytes.
Fda, Silver Spring, MD.
We studied the suitability of serum-free medium, reduced serum medium, and
chemically defined medium with serum replacement for growing rat neonatal car-
diomyocytes. The cardiac cells were grown for 48 hours under six different cul-
motion conditions: the base make up (Dulbecco's Modified Eagle Medium (DMEM)
containing 100 μM BEdU, 10 mM HEPES, 50 μM penicillin and
50 μg/mL streptomycin) with 10% Fetal Bovine Serum (FBS)-(A), or 10% Goat Serum-
(C), or 10% Knockout Serum Replacement (KSR)- (D) or no serum – (B);
Knockout DMEM(Gibco) (E) and Advanced DMEM(Gibco) (F) were supple-
mented by 15% KSR and 2% PBS, respectively. The beating rates and viabilities of the cells were evaluated by counting the beats of cells under a phase contrast microscope and Neutral Red Assay method, respectively. The cell damage and cytotoxicity were de-
termined by measuring lactate dehydrogenase (LDH) activity and the troponin I
release from the cardiomyocytes were examined by rat cardiac Troponin ELISA.
The results indicate poor cell beating and viability for medium C without serum. The
media D, E, and F had relatively low background troponin level, provided
good viability and morphology and low cytotoxicity, and may be used for the car-
diotoxicity study.

Understanding the Relationship of PI3K Inhibition to
HERG Liabilities and QT Prolongation.
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Groton, CT.
Off target promiscuity is one of the biggest issues that kinase targeting drug discov-
ery teams have to overcome in order to achieve efficacy with limited toxicity. In a
recent paper, Zhongju Lu et al (Science Translational Medicine 25 April 2012 Vol 4
Issue 131) showed that suppression of Phosphoinositide-3-Kinase(PI3K) signaling
directly or indirectly via tyrosine kinase inhibition prolongs the QT interval by af-
flicting multiple ion channels, including HERG. In an effort to understand the role
of PI3K inhibitors and HERG channels, we investigated the effects of Pfizer PI3Kα
inhibitors both in a Dofetilide binding assay that determines their affinity for
HERG channels as well as in a patch clamp assay that assesses their functional ac-
tivity on these channels. Out of 48 PI3Kα inhibitors selected from four structurally
distinct series and with PI3Kα Ki less than 1 nM, 47 exhibited Dofetilide Ki values
> 10 nM, and one demonstrated a Ki value of > 2.0 μM, a concentration > 2000-
fold above its kinase activity. 8 compounds, including the one with a Dofetilide Ki of
2.0 μM, were tested in the HERG functional assay and all showed IC50 values > 30
μM. In addition, 16 marketed drugs, which are known HERG blockers and that have
been shown to induce QTc prolongation (14/16) and/or cause Torsades de Points in
the clinic (8/14), were tested in a PI3Kα enzyme assay. Only Verapamil, a cal-
cium channel blocker that inhibits HERG current (IC50: 0.2 μM) but does not pro-
long the QTc interval or cause TdP in the clinic exhibited activity on PI3Kα
(IC50: 33 μM), while the rest showed IC50 > 100 μM.
In summary, the data obtained from both Pfizer potent PI3Ka inhibitors and mar-
keted drugs demonstrate the absence of the correlation between PI3Ka inhibition
and HERG activity. Furthermore, our results support the notion that HERG inhibition, in the absence of
PI3Kα inhibition, is sufficient to prolong the QT interval.

Characterization of Cell Signaling Events in Human
Cardiomyocytes.
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MD.
Cardiotoxicity is a significant concern for anticancer therapeutics. “On-target” tox-
icity may result when a drug target that regulates cancer growth also serves an im-
portant role in normal cardiac function. To explore mechanisms of cardiotoxicity

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Cardiotoxicity is the leading cause for late stage drug attrition and withdrawals from the market. Serious adverse cardiac events have emerged as a prevalent risk for kinase inhibitors (KI). Although current in vivo screens can reduce known risks due to arrhythmia, there is urgent need for novel in vitro assays with sufficient throughput to identify risks and support the development of SAR against other prevalent cardiovascular (CV) toxicities. Recently, cellular impedance technology has been adapted for detecting spontaneous, synchronous beating of cultures of cardiomyocytes (CM) in a real-time, label-free format. Impedance technology is a good candidate for detecting the pleiotropic cellular effects of kinases since it detects morphological changes thereby giving a readout that is downstream of key toxicity targets in the contraction cascade (i.e. cardiac action potential, calcium flux, mechanical elements of contraction). We evaluated the application of impedance-based assays for screening KI effects on rat neonatal CM. We selected compounds from a MAP-microtubule affinity-regulating kinase (MARK) inhibitor program that failed in late-stage preclinical development with dramatically decreased blood pressure in anesthetized dogs as an example for the earlier detection of CV toxicity. Two MARK inhibitors were tested for their dose-dependently induced changes on CM beat rate and amplitude without causing cell death as judged by cell index, cellular ATP levels, or cardiac troponin release assays. The relative potency of the two compounds on reducing beat amplitude in impedance assays (EC50 = 4.31 μM and 0.55 μM; 20 min exposure) aligned with the ~10-fold difference in affinity for MARK isoforms 1-4. Knockdown of pan-MARK expression reduced beat amplitude by 40%. These MARK data indicate that impedance assays can specifically detect non-cytotoxic functional effects of KIs on CM. Our data support the validation of cellular impedance-based assays for an early preclinical CV toxicity screening of KIs.

83 Activation of Human Monocytic NADPH Oxidase by Chlorinated Cyclodiene Insecticides.

L. C. Mangum, J. E. Chambers and M. K. Ross, CVM Basic Sciences, Mississippi State University, Mississippi State, MS.

Although the mechanistic relationship between bioaccumulative organochlorine (OC) insecticide exposure and increased atherosclerosis risk is poorly defined, elevated systemic oxidative stress stemming from OC-mediated induction of NADPH oxidase activity may play a significant role in disease development. Activation of phagocytic N2-containing NADPH oxidase can result in a rapid intracellular accumulation of superoxide-derived reactive oxygen species (ROS) that may be directly linked to the progression of atherosclerosis. This study measured the ability of two legacy OC compounds to induce Nox2-containing NADPH oxidase activity in vitro, in addition to providing evidence for a possible mechanism of action. Human THP-1 monocytes exposed to micromolar amounts (1-20 μM) of the cyclodiene OC insecticide trans-nonachlor and dieldrin exhibited increased levels of superoxide phosphorylation of the p47phox regulatory subunit of NADPH oxidase, a necessary process for enzyme assembly and translocation, with trans-nonachlor demonstrating several fold greater potency than dieldrin. OC treatment also induced elevated levels of intracellular ROS, as shown by 2′,7′-dichlorofluorescein-diacetate fluorescence assay, suggesting increased superoxide anion production. Pretreatment of monocytes with arachidonyl trifluoromethyl ketone, a specific inhibitor of cytosolic phospholipase A2, prior to cyclohexane treatment abrogated p47phox serine phosphorylation and blocked the induction of arachidonic acid and prostaglandin liberation, as determined by UPLC-ESI MS/MS, suggesting that this enzyme may play a crucial role in the induction of NADPH oxidase activity by cyclodiene via the modulation of intracellular arachidonic acid levels. The results suggest that trans-nonachlor and dieldrin are capable of altering intracellular ROS levels via an...
Drug induced cardiotoxicity can manifest itself through a variety of mechanisms including mitochondrial toxicity, apoptosis, oxidative stress and ion channel disturbance. However, cardiac hypertrophy is the most common safety finding. Here, we aimed at establishing an in vitro platform which could reliably characterize hypertrophic responses. Fifty drugs reported to cause hypertrophy in vivo were utilized in this study and consistent of a variety of drug classes such as anthracyclines, statins, kinase inhibitors and catecholamines. Compounds were characterized in H9c2 cells assessing the following endpoints: cell size, protein synthesis, fetal gene expression (ANP, BNP, MYH7), cell viability and reactive oxygen species (ROS). Our results indicate that unlike the traditional cardiac hypertrophy which is driven by ventricular wall stress and hormonal factors including angiotensin II, endothelin-1, and catecholamine, compound-induced cardiac hypertrophy (increase in protein and cell size) may be initiated by the same mechanisms that drive cytotoxicity. In contrast, fetal gene expression is different from the mechanisms that drive cell size and protein increase. Specifically, our results demonstrate that: 1) all protein synthesis-inducing compounds (23) caused significant cytotoxicity, 2) all compounds causing increase in cell size also induced protein synthesis; 3) both protein and cell size increase were compound-concentration dependent with protein increase occurring always ahead of cell size increase; 4) of the total of 50 drugs known to cause hypertrophy in vivo, 28 compounds induced fetal gene expression; 5) Oxidative stress was associated with the hypertrophic response (25%). In summary, our data suggest the initial trigger(s) of compound-induced hypertrophic response are multi-factorial; one of which shares the same mechanism that induces cytotoxicity. Not all compounds known to cause hypertrophy in vivo were accurately characterized by our in vitro approach. Whether stem cells and additional mechanistic parameters can improve our assay is subject to future studies.

Highly predictive in vitro assays suitable for high throughput screening (HTS) for potential cardiotoxicity are critical to drug safety testing. Adult human stem cell derived cardiomyocytes show promise for screening compounds during early drug development. We developed methods for measuring the impact of drug candidates on the beating rate of human iPS derived cardiomyocytes using fast kinetic fluorescence imaging. Cardiomyocyte contraction rate and pattern are characterized by monitoring changes in intracellular Ca2+ measured using calcium sensitive dyes. The assay was optimised for HTS and allows characterization of beating profiles by using multi-parameter analysis outputs such as beating rate, peak frequency and width, or waveform irregularities. The assay is suitable for assessment of short-term (minutes) and delayed (days) effects. Next, we tested known cardiotoxic compounds including alpha and beta blockers, hERG inhibitors, ion channel blockers, etc.; as well as control drugs. IC50 values showed a significant rank correlation with published values determined by other cardiotoxicity models as well as good concordance with reported human plasma Cmax values. The assay was further tested using commercially available cardiotoxicity library representing different classes of compounds including receptor antagonists, ion channel blockers, anti-cancer and anti-inflammatory drugs, and kinase inhibitors. The estimated balanced prediction accuracy of the assay was greater than 80%, and multi-parameter characterization of beating profiles allowed identification of specific patterns defining hERG or Na channel blockers. We conclude that this assay shows utility for screening compounds for potential to cause arrhythmic and non-arrhythmic cardiotoxicity.

HIV patients undergoing antiretroviral therapy exhibit an increased incidence of cardiovascular events and their associated diseases. Though HIV therapy generally involves a combination drug approach, nucleoside reverse transcriptase inhibitors (NRTI) are considered a backbone of this therapy. Prior studies in rodents suggested that NRTI promote HIV-associated endothelial dysfunction, an initiating factor in atherosclerosis. Further, this cellular dysfunction was associated with mitochondrial injury. In cultured endothelial cells, NRTI treatment increased mitochondrial ROS production, while decreasing ATP production and the activities of mitochondrial electron transport chain (ETC) complexes I-IV. While these effects were observed for acute treatment, in other studies, we noted that when the mitochondria were exposed to low doses of NRTI, damaged mitochondria were re-activated. In these studies, human aortic endothelial cells (HAEC) were exposed to chronic NRTI treatment. Mitochondrial ROS production, while decreasing ATP production and the activities of mitochondrial ETC complexes I-IV, showed a delayed (days) effect. Next, we tested known cardiotoxic compounds including receptor antagonists, ion channel blockers, anti-cancer and anti-inflammatory drugs, and kinase inhibitors. The estimated balanced prediction accuracy of the assay was greater than 80%, and multi-parameter characterization of beating profiles allowed identification of specific patterns defining hERG or Na channel blockers. We conclude that this assay shows utility for screening compounds for potential to cause arrhythmic and non-arrhythmic cardiotoxicity.
calcification caused by 2 mM Pi by 25% and 59% in VSCMs, respectively. A dose-re- sponse relationship of calcification inhibition with increasing concentrations of py-rophosphate (PiP) revealed that the IC50 shifted from 4.26 μM to 2.14 in the pres-ence of 5 mM NaF, therefore indicating that less PiP is necessary to prevent calcification when F is present in the calcification medium. Similar effect was ob-tained with phosphonofluoridic acid. Electron microscopy (TEM, SEM, ED and EDS) observations have revealed that the calcification deposits consist of spherolites of intergrowth particles that had an amorphous character, a low density, and a laminar particle shape, in the absence of F-. With increasing F-concentration, the deposits turn more dense and compacted, while the particle shape becomes clearly fibrillar. F is not incorporated in the de-posits, thus suggesting a modification of crystallization process and inhibition of nucleation, which could explain the prevention of calcification. This effect was similar both in living and dead (fixed) cells, which indicates an intrinsic effect of F- ions on the calcification process independent of cellular activity. Furthermore, F-decreases the Pi-induced expression of osteogens, most likely as the consequence of the reduced calcium phosphate deposits.

In conclusion, addition to drinking water can have a beneficial effect in prevent-ing vascular calcification originated during hyperphosphatemic conditions.

91 Vascular Effects Induced by 15nm Gold Nanoparticles in Isolated Rat Aortic Rings.

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Gold nanoparticles (AuNPs) have been used in biomedicine as therapeutic tool. However, their role on the vascular physiology, have not been fully studied. The purpose of the present work was to evaluate the effects of 15 nm AuNPs on the vas-culature, using a rat aortic rings model, pre-contracted with phenylephrine, a well known contractile agent. Adult male Sprague Dawley rats were sacrificed; aorta was excised, and maintained in an organ bath chamber with physiological solution. Each individual aorta rings were treated under isometric conditions, in presence and absence of endothelium (E), along with increasing concentrations (0.1-100 μg/mL) of AuNPs. AuNPs exerted a vasodilator effect independent on E, the relax-ation was exerted in the same magnitude that was found in presence of E. Relaxation induced by the Au-NPs were not dependent on nitric oxide (NO), since, L-Nitro-arginine methyl ester (L-NAME), an inhibitor of the NO produc-tion and a potent vasodilator mediator, did not block this effect. However, a pre-treatment with indomethacin, an inhibitor of prostanooids synthesis, blocked partial-ly the vasodilation induced by AuNPs, suggesting that prostanooids could be, at least in part, mediators of these actions. Further studies are underway to evaluate the mechanisms of action which are underlying the relaxation induced by AuNPs, and the implications that they could confer in the cardiovascular physiology.

92 Nano-Cerium Dioxide Exposure and Arteriolar Dysfunction: Exposure Route Dependency.

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Nano-cerium dioxide (CeO2) is being used or developed as a fuel catalyst (diesel), a protective drug (radiation and ischemia) and as a contrast agent (medical imaging). These diverse uses possess the potential for human exposures that are beyond the well studied pulmonary route. Our laboratory has assessed the arteriolar effect of nano-CeO2, 24hrs post-pulmonary exposure; however, other routes of exposure such as systemic and gastrointestinal have not been investigated. Therefore, our aim was to analyze the microvascular effects of nano-CeO2 via instillation, injection, and gavage, Sprague-Dawley rats were intratracheally instilled (100 μg), intra-venously injected (100 or 900μg), or gavaged (100 or 600μg) with nano-CeO2 sus-pended in Normosol (5% serum). 24hrs later the mesentery was harvested, and 4th or 5th order arterioles were dissected and prepared for isolated vessel experiments. Arteriolar reactivity was evaluated by determining endothelium-dependent (acyetyl-choline, 10-4-10-5M), and -independent dilatation (permeine NONOate, 10-4-10-3 M), vasoconstriction [phenylephrine (PE), 10-4-10-3M], and mechanotransduc-tion [myogenic responsiveness (0-10-5 mMgH2) and shear stress (0-30 μl/min)]. Endothelium-dependent and -independent dilatation for all three exposure routes (instillation: 30% and 50%, injection: 45% and 55% and gavage: 36% and 64% vs sham arterioles, respectively) was significantly impaired. Arteriolar responsiveness to PE or pressure was unaltered for the exposure routes. There was an impaired di-lation in response to flow for the rats injected or gavaged (53% and 35% vs sham, respectively). This arteriolar dysfunction was exposure route dependent because lit-tle to no arteriolar dysfunction was observed in rats injected or gavaged with 100μg of nano-CeO2, as compared in instilled rats. These results provide evidence that mi-crovascular dysfunction is present 24hrs after nano-CeO2 exposure and is depend-ent on the exposure route.

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Simulated Metabolism in the Langendorff Isolated Rat Heart: A Comparison of Novel and Traditional Dosing Techniques for 5-Fluorouracil and Doxorubicin.

J. B. Ross, K. A. Henderson, R. Borders, P. S. Hong, W. M. Black and B. M. Roche. Battelle, Columbus, OH.

There is a clear need for more predictive and translatable assays in early drug discovery. It is important to take a multi-scale approach to translate data to pre-clinical and clinical trials. The Langendorff isolated heart model provides a tool to assess cardiotoxicity in the whole organ in the absence of in-vivo variables. We used the Langendorff to assess both acute and latent cardiotoxicity of 5-fluorouracil (5-FU) and doxorubicin. Physiologically Based Pharmacokinetic (PBPK) models were developed to determine clinically relevant doses. A reverse extrapolation from human physiological and physiochemical parameters to rat was performed. The resulting pharmacokinetic (PK) curves led to some challenging dose considerations; both 5-FU and doxorubicin are rapidly cleared in humans and rats, never reaching a steady state. Other PK parameters such as Area Under the Curve (AUC) and the maximum concentration (Cmax) pose problems as well. AUC will accurately represent the total amount of drug over a given period of time; however, a single concentration derived from the AUC could potentially underestimate a biological response. In contrast, Cmax would greatly overestimate the total amount of drug, potentially leading to false toxicity. Computer generated dosing schemes were developed and characterized in an effort to fit the modeled PBPK curves. Initial concentrations (Cmax) were diluted with deionized water and perfused through the Langendorff system in the absence of a heart. Perfusate was collected at various time points, analyzed with HPLC, and concentration curves were generated for proof of concept. An acceptable fit was observed between the computer generated and the tested curves. This dosing technique was applied to an isolated rat heart; multiple end-points were collected and compared to previous data collected at constant perfusion concentrations. The novel dose schemes proved to be a better representation of in-vivo concentrations in our ex-vivo model.

Oxidative Stress Uncovers β2-Adrenergic Mediated Dilation to Curcumin Mimicked by Preventing Clathrin Endosome Formation.

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The nutraceutical, curcumin, has anti-oxidative properties, but also is a potent adrenergic receptor (AR) agonist, with an EC50 for dilation in the nM range (beta, bAR). Our goal was to determine how oxidative stress impacted vasoactive responses to curcumin. Intravital microscopy of small arterioles in the hamster cheek pouch was performed (N=10; pentobarbital 70mg/kg i.p.). Oxidative stress (OX) was induced by applying exogenous nitric oxide for 2 minutes in the tissue bath. Before OX, dilation to micropipette applied curcumin was dose dependent, with a maximum from 10-9 to 10-6M. After OX, dilation was eliminated in the nM range and higher range, and significantly left shifted to a narrow peak from 10-15 to 10-10M, which was inhibited by the b2-adrenergic receptor inverse agonist, carazolol (1nM). We next tested whether preventing clathrin endosome formation with dynasore altered dilation to curcumin. Dynasore (80uM) alone eliminated dilation to curcumin in the uM range, and uncovered a dilation in the pM range. Dynasore with OX further increased the potency, and increased the efficacy of dilation to curcumin over the range 10-15 to 10-10M. To understand the mechanism of this OX induced narrow potency range, we tested whether curcumin and the bAR were internalized and co-localized using FRET in HEK cells. Before OX, curcumin was rapidly internalized (seconds), and co-localized with bAR. After OX (1μM CoCl2), curcumin was internalized, but bAR remained at the plasma membrane. Thus, in cells, OX prevented internalization of curcumin with bAR. In situ, inhibition of clathrin endosome formation mimicked and enhanced the vasoactive responses after OX. (AHA 6655908T)

Diminished Oxygen Supply in Microvessels following Inflammatory Oxidative Stress.


Increased heterogeneity of flow accompanies many inflammatory states, including oxidative stress (OX) with toxicant exposure or thermal injury. Red blood cell (RBC) flux along nutrient flow paths is an indicator of oxygen delivery. Many inflammatory mediators result in a specifically located arteriole-venule RBC shunt pathway. Our goal was to directly measure oxy- vs deoxy-hemoglobin (HbO v HbR) to determine whether OX induced shunt paths resulted in heterogeneity of oxygen supply. A controlled thermal injury model revealed that hamster cheek pouch tissue localized OX so that unequivocal control vs OX spatial locations could be observed. Adult male hamsters (N=10, isoflurane) were prepared for intravitral microscopy of the cheek pouch tissue. Fluorescently labeled RBC flow markers confirmed shunt pathways after a thermocouple controlled 50 degree C 500um spot burn (same location at 5mm below the skin). A low power laser speckle contrast imaging was used (1-2h after burn) to determine HbO and HbR at 630 and 570nm at locations within 500um of the burn (near) vs. locations >1mm from the burn rim (far); values of changes in HbO and HbR were obtained in [Hb] mM. Total Hb in the large vessels was consistent with a local tube hematocrit of 5-40%, as is typically found using fluorescent RBC. HbO was 30% less in perfused large arterioles near vs far from the burn, and 24% less in perfused large venules near vs far from the burn. Within the terminal arteriolar networks far from the burn, HbO is 7% higher along the shunt pathway vs the non-shunt pathway, consistent with RBC flux values. Near the burn, HbO is 71% greater along the shunt pathway vs the non-shunt pathway, also consistent with RBC flux values. Comparing HbO along shunt pathways, HbO is decreased 52% near vs far from the burn rim. Thus, in this model of inflammation, RBC flux and HbO values together suggest that a diminished oxygen shunt pathway is initiated by inflammation and maintained for several hours after the inflammatory event. (NIH HL55492; AHA 6655908T)

Validation of the Amphibian Metamorphosis Assay for Potential Endocrine Disrupting Chemicals with Xenopus Laevis.


The new European Union Plant Protection Products Regulation (PPPR 1107/2009) identifies the need to consider whether a substance is a potential endocrine disrupter in aquatic non-target organisms and the current draft of the PPP data requirements refers to three screening assays for ecotoxicological endocrine-disrupting potential. Of these, we describe in detail our experience in the establishment and validation of the amphibian metamorphosis assay (OECD 231; OPPTS 890.1100) with the African Clawed Frog. In this method, in order to satisfy validity criteria in the rearing phase, conditions necessary to allow tadpoles to develop from fertilisation to development stage 51 as defined by Nieuwkoop and Faber (1994) were established, and individuals selected for the exposure phase and transferred to test vessels. To establish the assay, 400 tadpoles were exposed to a range of levels of the three reference substances, thyroxine (T4) which produces stimulatory effects on the normal function of the hypothalamic-pituitary-thyroid (HPT) axis, sodium perchlorate (which retards development) and ionopanic acid (which affects hind limb development) and levels of each were verified using an appropriate analytical method. At Day 7, 80 randomly selected individuals at each exposure level were removed and assessed (body weight, development stage, hind limb and snout to vent length). Exposure continued for a further two weeks and the study terminated on Day 21 when all the remaining individuals were assessed as on Day 7. Following developmental stage matching, 80 individuals were selected for thyroid removal and histopathological analysis. We found that our test and the ring test results published by the OECD (Series on Testing and Assessment Document Number 77) and make a number of observations on methodology that may improve the reproducibility of these assays.

Mammary Gland Morphology and Gene Expression Signature of Prepubertal Male and Female Rats following Exposure to Exogenous Estradiol.

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In order to properly understand whether xenestrogens act as true estrogen agonists, it is essential to possess a solid portrait of the physiological effects of exogenous 17β-estradiol (E2). Because the estrogen-dependent gene expression is one of the primary indicators of estrogenic action, we have assessed effects of three doses of exogenous E2 (0.1, 1.0 and 10 μg/kg of body weight/day) on the mammary gland morphology and gene expression profiles of prepubertal male and female rats of both sexes compared to untreated controls. The mammary gland was more responsive to E2 treatment in males than in females with 1392 genes modulated >1.5-fold up or down relative to controls at the highest E2 dose compared to 463 genes. There

96 Validation of the Amphibian Metamorphosis Assay for Potential Endocrine Disrupting Chemicals with Xenopus Laevis.

94 Oxidative Stress Uncovers β2-Adrenergic Mediated Dilation to Curcumin Mimicked by Preventing Clathrin Endosome Formation.

93 Simulated Metabolism in the Langendorff Isolated Rat Heart: A Comparison of Novel and Traditional Dosing Techniques for 5-Fluorouracil and Doxorubicin.

95 Diminished Oxygen Supply in Microvessels following Inflammatory Oxidative Stress.
was an increase in the number of terminal end buds in males (P<0.05), and a corresponding increase in the expression of the gene encoding amphiregulin (P<0.05). A protein known to drive the differentiation of terminal end buds. In intact females, the highest dose of E2 tested induced an increase in the expression of genes encoding milk components, as well as muscle proteins. Lower doses had limited effects on gene expression. Therefore, the prepubertal rat male mammary gland is a very sensitive tissue to evaluate estrogenicity using morphological changes coupled to microarray-analysis of gene expression. Intact prepubertal females were comparatively poorly responsive to exogenous E2, although many modulated genes were common to both sexes. Supported by USDA CRIS-6251-51999-007-04S.

Potential Endocrine Disruption of a Drinking Water Sample from the State of São Paulo, Brazil.

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Many xenosterogens, natural and synthetic estrogens may end up in the water bodies through sewage discharges. Contaminated rivers are conventionally treated by Water Treatment Plants (WTP) to produce drinking water, but emerging contaminants may remain. This study investigated the potential for endocrine disruption of a drinking water sample from a São Paulo State WTP, using two harmonized biosays. Female rats 21 days old were exposed by gavage to drinking water extracts during 03 days (uterotrophic assay; OECD 440) or 20 days (pubertal development female rat assay, EPA 890.1450) at 33.3, 166.5, and 333.0 mL equivalent of water/kg body weight, modeling a daily ingestion of 2 L, 5 L and 10 L of drinking water by a 60 kg human being. Traces of caffeine (5.8 ug/L), estrone (1 ng/L) and atrazine (11.2 ng/L) were detected in the extracts by LC-MS/MS. Androgen or estrogen agonistic activity of the water extract resulted negative by the bioluminescent yeast assay (BLYES, BLYAS). Accordingly, both in vivo assays were negative: there was no significant increase of the uterus wet weight in the uterotrophic assay and no alteration of the moment of vaginal opening in the pubertal assay. However, the serum levels of LH, FSH, PRL and 17β-estradiol were altered, with a significant dose-response increase. Therefore, it has to be assumed that the tested drinking water sample induced steroidogen dysregulation in vivo.

Development of Medium-Throughput Thyroperoxidase Inhibition Assays for Screening.


Thyroperoxidase (TPO), the catalyst for thyroid hormone (TH) synthesis, is a target for thyroid-disruptors, including methimazole (MMI), isoflavones, benzoquinone-2, and malachite green; however, no medium-to-high-throughput screening methods for TPO inhibition are available. To adapt the low-throughput guaiacol oxidation assay for TPO inhibition to chemical screening, we replaced guaiacol with a fluorescent peroxidase substrate in a rat thyroid-based assay and developed an in vitro human TPO model. We tested the hypothesis that use of a peroxidase substrate (Amplex UltraRed, AUR, LifeTech) in 96- or 384-well plate formats with automated reagent delivery could increase the TPO assay throughput with thyroid microsomes. The IC50 of MMI-induced TPO inhibition was reduced with AUR compared to the guaiacol assay, with average IC50s of 0.15 μM and 5 μM, respectively. AUR signal was stable from 30-120 min after initiation. The dynamic range of the assay with MMI was 6- to 10-fold using 96- or 384-well formats, with Z’ scores of 0.75 to 0.9. A 21 chemical training set is being used to validate the assay for screening. One limitation of this model is the availability of human thyroid microsomes. Preliminary work demonstrated that lentiviral-transduction of a human cell line (HEK293T) with recombinant TPO could also be used to test for human TPO inhibition. TPO activity measured by the AUR assay in cell fractions was maximally inhibited by MMI at 100 μM. This approach to improving and developing medium-throughput assays for thyroid-disruptor screening demonstrates the feasibility of screening 1000s to 1000s of chemicals for TPO inhibition, and drastically reduces the need for animal tissue. This abstract does not necessarily reflect the policy of the US EPA.


Substantial public health concerns exist over the potential endocrine disrupting capabilities of a wide variety of untested or under-tested natural and industrial chemicals. It is clear that the development of accurate, high-throughput, and inexpensive testing regimes will be key to mitigating public concern. Here we report on the development of a novel screening assay for estrogenic activity that utilizes an autonomously bioluminescent human cell line to provide direct bioavailability data. To construct this cell line, estrogen-responsive human breast carcinoma cells (T-47D) were genetically engineered to express the full bacterial bioluminescence gene cassette (luxCDABE;frp), generating an autonomously bioluminescent cell line (T-47D/Lux) capable of maintaining bioluminescence output independent of substrate addition. Bioluminescence emitted from T-47D/Lux cells was correlated tightly (R2 > 0.99) to the number of cells present in a population, permitting the use of light production dynamics as an indicator of cell proliferation. Additionally, the substrate-free nature of the lux system allowed for continuous, near real-time monitoring of the same cell population throughout exposure to the tested compounds. A significant change in bioluminescence production (p < 0.05) compared with unexposed control was observed 3 days after exposure to concentrations of 17β-estradiol (E2) as low as 1 μM. The EC50 for E2 in this assay was determined to be approximately 10 μM. These results are similar to those obtained using a traditional cell proliferation assay, but offer the advantage that data acquisition can be performed in a fully automated fashion since the need for sample extraction or substrate addition is removed, making it an ideal candidate for high-throughput analysis.

Cryopreserved Rainbow Trout Hepatocytes Model Endocrine Disruption As Detected by Protein and Gene Expression of Vitellogenin.

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The toxicity of environmental pollutants may occur through modes of action that alter normal estrogenic functions, resulting in altered development, growth and reproduction of aquatic species. Consequently, there is a need to develop assay endpoints capable of identifying endocrine disruption in fish. Our lab has previously shown the utility of cryopreserved trout hepatocytes. Here we use this cell culture method to identify the potential estrogenicity of chemicals through vitellogenin (VTG) expression. This biomarker indicates an estrogenic effect when the protein is expressed by male trout hepatocytes, both in vivo in plasma and in vitro in cell culture media. To test the performance of cryopreserved trout hepatocytes, reference chemicals estradiol (E2), estrone (E1) and diethylstilbestrol (DES) were used and culture media was collected for ELISA 96-hrs post-treatment. Our results show that the levels of protein detected with cryopreserved trout hepatocytes are similar to results previously published using fresh trout hepatocytes, ranking the compounds E2 ≥ DES > E1. To determine whether transcriptional regulation of VTG correlated with protein expression, quantitative real-time PCR was performed. mRNA was collected from cryopreserved hepatocytes treated for 24 hours with the same doses of reference chemicals. VTG gene expression correlated well with protein expression,-ranking the compounds similarly. In addition, the negative control corticosterone showed no increase in gene expression at any dose, indicating that this is a sensitive, yet specific endpoint. These studies demonstrate both ELISA and transcription-based methods can be used with cryopreserved trout hepatocytes for screening chemicals for potential estrogenic effects, enabling greater access to this model system and potentially improving the turn-around time for generating results through detection of VTG mRNA levels.

Permethrin Does Not Have Endocrine Disrupting Properties As Evaluated by the US EPA's Endocrine Disruptor Screening Program (EDSP) In Vito Assays.

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Permethrin, a pyrethroid insecticide, was evaluated for potential endocrine activity in 3 of 9 in vivo US EPA EDSP assays. The maximum tolerated dose (MTD) of permethrin used in these assays was selected based on the results of dose range-finding studies. Sprague Dawley rats were administered permethrin in corn oil by oral gavage for 10 (Hershberger Bioassay; OPPTS 890.1400, castrated model), 21/22
clearance appears evident. Similarly, results for thyroid effects in the pubertal assays were consistent with MoA assigned an indirect MOA due to enhanced thyroxine clearance not relevant to humans. Furthermore, benefit was negative in the amphibian metamorphosis assay designed to detect thyroid agonists and antagonists. A high-degree of coherence and consistency is observed in results from the higher-tiered data and previous toxicity studies showing no direct interaction with the endocrine system. In conclusion, the vast majority of Tier 1 studies/endpoints were clearly negative and those few positive responses were seen only at high-dose levels and were either isolated (e.g., ERTA) in nature or secondary to other effects.

105 Endocrine Exposure at Environmentally-Relevant Concentrations.

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Endocrine disruption has become an important topic of public concern. Despite an increasing amount of attention, little is understood about how doses of endocrine disrupting chemicals (EDCs) at environmental concentrations affect homeostasis. To address these concerns, we performed a pre-/post-natal reproductive toxicity study to measure the developmental toxicity of low doses of three anti-androgenic compounds: vinclozolin, flutamide, and prochloraz. The tested doses were selected to mimic low-effect levels, the no observed adverse effect levels (NOAEL) for endocrine effects, and the acceptable daily intake (ADI). Despite mild maternal toxicity in parental females treated with the top dose of prochloraz, sufficient offspring were produced to evaluate the developmental effects of the three EDCs. While female offspring developed normally, the male offspring showed effects known for anti-androgens. One sensitive clinical parameter was the retention of nipples and/or areolas by male animals on PND 12. This effect was largely transient, as all had regressed by PND 21, with the exception of the offspring exposed to the flutamide top dose. The young adult male offspring displayed additional anti-androgenic effects including, delayed sexual maturation and reduced male sex organ weights. These alterations were observed at the effect doses of all three compounds, and at the putative flutamide NOAEL. Offspring from the flutamide top dose group had an increased incidence of developmental sexual defects including hypospadias, short penis, and cryptorchidism. No effects at all were noted at the low doses, as expected for the ADI. Assessment of sexual steroid hormones and their precursors revealed no effects at any of the dose levels. Taken together, the weight of evidence of the clinical and pathological findings suggests a lack of a non-monotonic dose-response curve.

106 Assessment of Estrogenic and Androgenic Impairment of Iowa Surface Water by Chemical Analysis and Bioassays.

S. Eto1, J. Vargo2 and G. Ludewig2. 1University of Iowa, Iowa City, IA; 2Iowa Hygienic Lab, Coralville, IA.

Limited data are available regarding the presence of hormones and steroids in Iowa surface water. To obtain more information about the hormonal impairment, river water samples were collected in October 2011 in two different locations in the Iowa City and Des Moines. Chemical analysis of water extracts was performed by LCMSMS using drinking water method EPA 539 that assesses equilin, estriol, 17α- and 17β- estradiol, estrone, 17α-ethynylestradiol, androstenedione, testosterone, progesterone, and 17β-trenbolone concentrations. For endocrine disrupting activity of the water extracts the E- and A-screen with human MCF-7 breast cancer cells (estrogenic and androgenic activity) and the H295R assay with adrenal cancer cells (interference with steroid hormone synthesis) were used. A goal of this study was to determine how well chemical testing for targeted hormones and steroids reflects the total androgenic and estrogenic activity determined by bioassay tests. Preliminary evaluation revealed that the added preservative 2-mercaptopyridine-1-oxide or a dehydrating agent from the flutamide top dose group had an increased incidence of developmental sexual defects including hypospadias, short penis, and cryptorchidism. No effects at all were noted at the low doses, as expected for the ADI. Assessment of sexual steroid hormones and their precursors revealed no effects at any of the dose levels. Taken together, the weight of evidence of the clinical and pathological findings suggests a lack of a non-monotonic dose-response curve.

103 Linking In Vivo Enzyme Activity To In Vivo Effects: Thyroperoxidase Inhibition in Methimazole-Exposed Rats.


The anti-thyroid drug methimazole (MMI) inhibits thyroperoxidase (TPO), the enzymatic catalyst for thyroid hormone (TH) production and a target for thyroid-disrupting chemicals. Our development of a medium-throughput screening assay for TPO inhibitors using a fluorescent peroxidase substrate (Amplex UltraRed) has instigated characterization of the relationship between in vitro TPO inhibition using thyroid microsomes and in vivo TH status. This work tested the hypothesis that TPO activity would be decreased in thyroid from rats exposed to MMI that decreased serum THs. Weanling female Long-Evans rats received MMI in corn oil (0, 0.1, 0.3, 1, 10, 20, 30 mg/kg/day po) for 4 days and were sacrificed 24 hr later. Pooled thyroid microsomes (n=8/pooled treatment) were used in the TPO inhibition assay. As a positive control, the vehicle control microsomes were exposed in vitro to MMI (0.015 - 200 μM), which yielded an assay dynamic range of 6-fold, an IC50 = 0.14 μM, and z’ = 0.9. TPO inhibition was in a dose-dependent manner, up to 53% of vehicle control at 30 mg/kg/day, which corresponded to significant serum thyroxine (T4) decreases of 60% and thyroid gland weight increases of 64% (p<0.05). TPO inhibition in the 10-30 mg/kg/day MMI corresponded to in vivo MMI concentrations of 0.14-0.44 μM. These results suggest that MMI-induced serum T4 decreases in vivo correspond to significant TPO activity decreases in vitro. Continued work is aimed to strengthen understanding of the association between in vitro TH perturbation and dynamics of TPO inhibition in the AUR screening assay. This abstract does not necessarily reflect the policy of the US EPA.

104 Endocrine Disruption Potential of Benefin: Weight of Evidence Evaluation.

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Benefin, a pre-emergent herbicide, was included on List 1 of chemicals to be screened under US EPA’s Endocrine Disruptor Screening Program (EDSP). Following completion of 11 Tier 1 EDSP assays, a Weight of Evidence (wOE) evaluation was conducted using all available data to determine if benefit has potential to interact with the estrogen, androgen or thyroid (EAT) systems following the five (5) – tiered OECD conceptual framework. For the in vitro assays, benefit was negative for androgen and estrogen receptor binding, and negative in the aromatase and steroidogenesis assays but induced a weak positive response in the estrogen receptor transactivation assay (ERTA). However, this slight positive response in the ERTA assay was not corroborated in other estrogen sensitive assays estrogen receptor (ER) binding, uterotropic, estrogen-sensitive endpoints in the female pubertal, and vitellogenin levels in the fish short-term reproduction assay (FSTRA) that were negative. Anti-androgenic and anti-estrogenic effects were observed in the pubertal assays and in the H295R assay only at the high dose levels. However, the wOE evaluation of the data does not support a direct interaction; rather, an indirect mode-of-action (MoA) due to enhanced liver enzyme induction and hormone

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Endocrine Disruption: Weight of the Evidence for Low-Dose Effects of TCDD on Sperm Counts.


Considerable debate is ongoing regarding the potential for endocrine-mediated low-dose effects. One of the examples that has been presented as evidence for low-dose effects is exposure to 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) and the impact on sperm counts in the male offspring. Although one epidemiological study has reported reduced sperm counts in males exposed to TCDD, the results of this study are limited by the number of subjects, evaluation of a single sperm sample, potential differences in sexual activity, and lack of control in the collection/trans- port of the sperm sample. We have conducted a weight-of-the-evidence review of the toxicological studies investigating the impact of maternal TCDD exposure on sperm counts in their offspring and the results have been highly variable. Epididymal sperm counts have been reported to be reduced at doses as low as 64 ng TCDD/kg, but other studies have failed to show effects at doses of 800 ng TCDD/kg or higher. Several other animal studies have not reported effects on sperm counts at doses between 10 and 50 ng TCDD/kg, suggesting a potential threshold. While sperm effects have been reported in experimental animal studies at 64 ng TCDD/kg, this dose should not be considered a low dose because human exposures are much lower. This dose is not relevant to current human exposure to TCDD based on estimated intakes that are 100-fold lower for all dioxins and furans combined based on TEQ (toxic equivalents). Thus, discussions regarding the potential endocrine disruptive effects of TCDD on sperm counts are misleading and do not occur at the low doses experienced by general American public.

Effects of Pesticides on Cytochrome P450 17 (CYP17) and Androgen Receptor (AR) Function in Human H295R Adrenocortical and LNCaP Prostate Cancer Cells.

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Exposures to endocrine disrupting chemicals, including pesticides, are thought to be contributing to the increased incidence of certain endocrine-related cancers in the human population, such as prostate cancer. Prostate cancer growth is initially androgen-dependent and under control of the nuclear androgen receptor (AR), which increases the gene transcription of proteins involved in cell proliferation, such as prostate specific antigen (PSA). CYP17 is a key enzyme in the biosynthesis of androgens and increased expression is associated with increased prostate cancer risk. We evaluated the effects of several pesticides suspected or known to modulate hormonal function in androgen-dependent LNCaP human prostate cancer cells and in an in vitro steroidogenesis model, the H295R human adrenocortical carcinoma cell line. Benomyl, vinclozolin and prochloraz reduced dithydropyrostero- steron (DHT)-stimulated LNCaP cell proliferation and nuclear AR protein accumulation concentration-dependently (0.1–30 μM). Levels of an active phosphorylated form of AR, pAR-Ser81, were increased by 10 nM DHT, whereas benomyl, vinclozolin and prochloraz decreased these stimulated levels (after a 1 or 6 h exposure). All three pesticides reduced DHT-stimulated PSA secretion by LNCaP cells. We found AR to be expressed in H295R cells but levels of AR and pAR-Ser81 were not affected by DHT, although they were increased by 30 μM atrazine. Benomyl and vinclozolin (10 and 30 μM) and prochloraz (1 and 3 μM) decreased CYP17 mRNA expression in H295R cells at sub-cytotoxic concentrations and prochloraz strongly inhibited CYP17-catalyzed conversion of pregnenolone to DHEA (24h exposure). In H295R and LNCaP cells, CYP17 protein expression was increased by atrazine (30 μM). In conclusion, certain pesticides exert combined antiandrogenic effects in androgen-dependent prostate cancer cells at the level of CYP17 and AR, the latter by reducing AR phosphorylation at serine 81.

Metabolomics Characterization of the Effects of Estradiol in Human Breast Cancer Cell Lines.

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Estradiol (E2 or 17β-estradiol) is the most potent naturally occurring estrogen. Its effects on cell function during development and in adults in a variety of tissues are mediated through estrogen receptors located in the nucleus, cytoplasm, mitochondria and at the cell membrane. Our objective is to employ high-throughput liquid chromatography mass spectrometry based metabolomics analysis to map pathways affected by estradiol in MCF-7 and T47D cells and then compare them with pathways affected by endocrine disrupting compounds (e.g. BPA and others). Concurrent cell growth and gene expression studies are also being carried out. Cells were exposed to estradiol at different concentrations and times. Stimulation of cell growth was observed at 0.1 nM in association with changes in the metabolome at 6 and 24hrs. A time course study at 100 nM estradiol revealed that some metabolites were significantly altered at various time points. The metabolites and pathways most affected included metabolites from cellular energy-related pathways (e.g. Malate- Fumarate shunt and alpha-ketoglutarate synthesis pathways (e.g. Arginine, Valine, Ornithine, Leucine/Isoleucine, 2-Methylmalate). Further investigation is underway and should give more insight into the identification of metabolism pathways affected by estradiol and endocrine disruptors.

Creating In Vitro Test Methods for Endocrine Disruptor Risk Assessment: Assay Development.


Endocrine disruptor test programs in the US are moving forward to identify active compounds and prioritize them for subsequent in-life toxicity studies. Our focus is on developing in vitro tests for cellular responses to endocrine active substances that will be sufficient for health risk assessment without moving on to in-life toxicity tests. We have begun a research effort for the estrogen receptor (ER) pathway to: 1) map signaling pathways for estrogen mediated proliferation, 2) define the dose-response for stimulation of estrogen signaling by xenobiotics and 3) develop computational models to predict chemical effect on uterine epithelium. The first phase develops a cell-based assay in human Ishikawa endometrial cell line expressing the three major ERs: ER1, ER2 and G-protein-coupled receptor GPER. The dose-response for 17β-estradiol (E2) and 17α-ethynyl estradiol (EE) (0.001 – 10 nM) were examined at 1, 2, 3, 4, 5, and 6 days of exposure for proliferation and induction of protein and gene targets of ER1, including alkaline phosphatase (ALP), proliferation associated gene GBE1 and progesterone receptor (PGR). E2 and EE increased proliferation and ALP activity by day 3 at 0.01 nM, while gene induction (GBE1, ALP, PGR) occurred earlier (day 1). Cells were also tested with selective receptor agonists PPT, DPN, and G1 targeting ER1, ER2 and GPER. PPT increased proliferation, ALP activity and gene expression similar to EE. DPN induced proliferation and ALP only at higher doses associated with cross-reactivity with ER1. GPER plays a role in regulating uterine proliferative response; however, we saw no change in proliferation upon treatment with G1 alone. This may result from low GPER expression in our cells. We also conducted expression analysis of studies with Ishikawa cells from our laboratory and published sources to evaluate dose-response of GO-categories by these receptor specific ligands. In vitro responses have been analyzed in light of in vivo rodent uterotrophic assay transcriptomic data to show consistency between in vitro and in vivo assays.

Leaching of Chemicals with Estrogenic Activity from Packaging into Popular Lab Animal Feeds.

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A robotized MCF-7 cell proliferation assay currently undergoing validation by IC-CVM/NICEATM was used to quantify the total estrogenic activity (EA) of chemical mixtures leaching from lab animal feed bags into feed over time. A standard open-formula (low phytosterogen AIN-93G) feed was purchased from Harlan®, Research Diets™ and TestDiet®. The TestDiet® feed was packaged in EA-free bags recently developed by PlastiPure for PMI® LabDiet® and other customers. All diets were analyzed 0, 2, and 4 weeks after purchase. All leachates were quanti- fied relative to the percentage of the maximum DNA produced per well induced by a test chemical with respect to the maximum DNA produced per well induced by the 17β-estradiol positive control (%RME2) and corrected by the cell response to the vehicle (negative) control. The significance level (p < 0.01) to detect EA was a %RME2 value 3 standard deviations more than the VC. According to this con- servative criterion, EA levels in 72 hour ethanol extracts of commercial bags for TestDiet®, Harlan®, and Research Diets™ were non-detectable (ND), signifi- cantly positive, and significantly positive, respectively. In the feed study, there was no detectable EA activity in TestDiet® feed at any time point. Feed samples from Harlan® and Research Diets™ were ND at week 0 but significantly positive following storage for 2 and 4 weeks after purchase. EA readings for Harlan® and Research Diets™ increased from week 0 to week 2 and again from week 2 to week 4. These data suggest that leaching of chemicals having EA from the non-EA free plastic packaging can alter the activity of the non-purified animal feeds. Such leaching of chemicals having EA from plastic packaging into animal feeds may impact research protocols that examine hormonally- responsive end-points.
The use of Novel Assays to Screen Large Chemical Inventories for Functional Estrogen Receptor (ER) Activity.

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Rapid, efficient screening assays are important for providing the means to prioritize large inventories of environmental chemicals for potential endocrine disrupting effects. Past efforts focused on transcriptional activation (TA) assays for estrogen, androgen, or thyroid receptors. The complexities of nuclear receptor (NR) biology, however, limit this approach as a comprehensive strategy to identify all NR modulators. For example, selective ER modulators can have cell type-specific activity dependent on the particular coregulator proteins expressed. We thus examined alternative, complimentary assay methodologies that provide unique functional information for chemical modulation of the ER pathway. We measured the ability of 1848 chemicals to induce ER dimerization and nuclear translocation, initial and required steps in ligand-induced activity of the ER pathway using the fluorescent imaging in intact cells at multiple time points. We also determined receptor selectivity using distinctly tagged ERα and ERβ receptors and found that the great majority of compounds to have similar potency for both receptors. Secondly, we determined the activity of these 1848 chemicals in an ER/chromatin assay that visualizes active ER transcriptional loci and provides functional information as to the level of agonist versus antagonist activity of the ligand. Overall assay showed chemicals could be categorized into several groups. First, there were unequivocal ER agonists active under virtually all testing conditions. Second, antagonists and partial agonists with weak or no activity in TA assays run in agonist mode were readily detected by these assays. Finally, there are spurious or artificial results for each assay that must be taken in to account when performing large scale assays such as this and are apparent by looking at data across multiple assays and formats. These results demonstrate the value of multiple, complimentary assay approaches in understanding complex biological responses with in vitro systems.

Effect of Technical Variation and Bioinformatics on the Biological Discovery in a Mechanistic Toxicogenomics Study.

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Whole-transcriptome sequencing using next-generation sequencing technologies, i.e., RNA-Seq, has drawn a significant attention as a groundbreaking tool for clinical application and safety assessment. However, critical assessment of RNA-seq to toxicology needs to be carefully conducted to understand whether the technology is robust and reproducible and how the choice of the bioinformatics approaches impacts study of the toxicity mechanisms and predictive toxicology. In this study, we conducted a specific study design to evaluate the impact of sequencing platforms and library preparation to the differentially expressed genes (DEGs) obtained, as well as the impact of different methods for DEG identification. Specifically, RNA samples from six rats’ livers were collected, three of them were treated with aflatoxin (AFL) and the other three were matched controls. Two libraries were separately prepared for the samples and two sequencing platforms (Illumina HiScanSQ and HiSeq2000 systems) were used. The investigation was based on the results derived from 6 different bioinformatics pipelines; each pipeline consists of a workflow including a specific genome trimming and mapping algorithm, quantification, and normalization. Several methods for DEG identification were compared. The preliminary results show that library preparation introduces more variance than sequencing platform on gene expression quantification. Additionally, more variance has been observed for the low-expressed genes and thus increases the false positive rate in DEG identification among those genes. We proposed two filters for data reliability control prior to DEG analysis, one on the low expression level and another on high variance. Comparison was also carried out at functional module and pathway levels.

Structural Classification of 1848 Chemicals Evaluated for Estrogenic Activity in 13 HTS Assays.

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High-throughput screening (HTS) assays are seeing increasing use for identifying chemicals that can cause toxicity via key biological pathways. However, biological complexities often mean that a single assay will fail to correctly classify all compounds for pathway activity. Here we use multiple orthogonal HTS assays to classify chemicals for their ability to interact with the estrogen receptor (ER) pathway among a structurally diverse library of 1848 chemicals, including pesticidal actives and inert, industrial chemicals, food additives, cosmetics and drugs. The assays are: reporter gene assays in HepG2 cells selected to maximize metabolic activity, HEK293 cells, BG1 cells, CHO-K1 cells; cell-free binding in HEK293 cells; a transcription-factor/DNA binding assay in HeLa cells; and an ER-sensitive cell proliferation assay in T47D cells. Multiple assay readouts used (fluorescence,
luminescence, sequencing, electronic impedance) allowed determination of possi- ble assay performance. New in vitro assays were mainly based on gene expression (cytofluorescence, unrelated endpoints) in the same engineered cells were also included. Assay data were compared with chemical structure clusters. A total of 237/1848 chemicals were active in at least 4 assays. (Note: the library was enriched in chemical categories expected to be estrogenic). This survey identified well known structural classes of xenobiotic compounds including steroids, phenols, chlorobenzenes, anilines and perfluorinated alkanes. In metabolically competent assays, we also observed bioactivation in classes of compounds including perethynils and antrachine derivatives. In addition, we confirmed classes of compounds that consistently cause false activity in certain assays, including surfactants. Finally, there were novel compounds active in many assays that will require more analysis and follow-up studies to confirm their activity. This abstract does not necessarily reflect Agency policy.

The FDA’s Center for Tobacco Products (CTP) has received a large volume of documents from its regulated industries, and even more documents are forthcoming. The documents are unstructured and in various formats, including email, PowerPoint slides, memoranda and others. These documents contain diverse information related to such areas as marketing, safety and risk. The language is free text often containing ambiguous semantic descriptions, posing difficulty in retrieving useful information in a consistent and accurate fashion that is needed for regulatory review. Moreover, the sheer volume makes manual reading of the entire corpus impractical. To increase the review efficiency, the FDA requires submitters to include key word tags corresponding to FDA supplied question areas. The goal of this study is to identify the hidden topics or concepts from the submitted documents using topic modeling, and to assess whether the extracted topics reflect the FDA defined questions and thus facilitate review. As a pilot study, more than three thousand thousand related tobacco documents were used to develop a topic model with latent Dirichlet allocation (LDA). The number of topics was optimized based on an information-loss approach. After generating the topics, a cosine similarity based approach was used to map FDA questions to the extracted topics. The results demonstrated that the extracted topics reflect the concepts of regulatory questions. The majority of documents clustered by topic modeling coincided with those grouped by FDA defined topics. This study demonstrates the potential utility of topic modeling as an unsupervised machine learning technique to classify the submitted documents based on the hidden topics, and thus provides an alternative means to support the review process.

The appearance of skin tumors after chronic dermal administration, which has cost and time constraints. We wish to identify biomarkers for petroleum-induced skin carcinogenicity. TCS (2,4,4′-trichloro-2′-hydroxydiphenylether) is a broad-spectrum antimicrobial agent used in personal care products. It has cytotoxic and endocrine-disrupting properties, and may be bioaccumulated from daily use. However, its impact on human health is largely unexplored with little evidence for deleterious effects in humans. A total of 248 proteins involved in different pathological processes were evaluated by in silico virtual screening to find new possible human protein targets for TCS. The 3D-structures of the proteins were downloaded from PDB, prepared and optimized by Sybyl X-2.0. The TCS structure was optimized by DFT at the B3LYP/6-31G level in Gaussian 0.9. Docking studies were performed in AutoDockVina 2.0 using a blind docking strategy; exhaustiveness of 25 and 10 runs. Calculated binding affinities were then employed for ranking proteins. A re-docking step of 100 runs and exhaustiveness of 100, as well as the interaction analysis with Ligand Scout, LigPlot and MMV were carried out on the complexes presenting the best docking results. The greatest affinity scores were found for CDC2-like kinase 4 (CLK4, -8.4±0.0 kcal/mol), oligomeric death domain complex (CRADD/PIDD, -8.3±0.1 kcal/mol); progestogen receptor (PGR, -8.1±0.1 kcal/mol) and estradiol 17-beta-dehydrogenase 1 (17-beta-HSD 1, -8.1±0.1 kcal/mol). Proteins involved in endocrine disruption and breast cancer-related processes. Validation showed our protocol predicted the binding site for the 17-beta-HSD and PGR native ligands, in agreement with crystallographic data. Moreover, TCS interacts in the same binding site occupied by the 17-beta-HSD 1 native ligand, but in a different one from that used by progesterone in PGR. Protein-ligand binding involves hydrophobic and hydrogen bonding interactions. These results suggest TCS may target different proteins, probably altering pathways involved in pathophysiological disorders. Vice-Rectory for Research.


Gene Expression Analysis to Predict the Dermal Carcinogenic Potential of Petroleum Streams. K. Mathijss1, J. van Delft1, M. van Steensel2, K. Goyak1, H. Ketelbesrens1, L. Freeman1 and R. Phillips3. 1Department of Toxicogenomics, Maastricht University, Maastricht, Netherlands; 2Department of Dermatology, Maastricht University, Maastricht, Netherlands; 3ExxonMobil Biomedical Sciences Inc., Annadale, NJ.

Triclosan (TCS; 2,4,4′-trichloro-2′-hydroxydiphenylether) is a broad-spectrum antimicrobial agent used in personal care products. It has cytotoxic and endocrine-disrupting properties, and may be bioaccumulated from daily use. However, its impact on human health is largely unexplored with little evidence for deleterious effects in humans. A total of 248 proteins involved in different pathological processes were evaluated by in silico virtual screening to find new possible human protein targets for TCS. The 3D-structures of the proteins were downloaded from PDB, prepared and optimized by Sybyl X-2.0. The TCS structure was optimized by DFT at the B3LYP/6-31G level in Gaussian 0.9. Docking studies were performed in AutoDockVina 2.0 using a blind docking strategy; exhaustiveness of 25 and 10 runs. Calculated binding affinities were then employed for ranking proteins. A re-docking step of 100 runs and exhaustiveness of 100, as well as the interaction analysis with Ligand Scout, LigPlot and MMV were carried out on the complexes presenting the best docking results. The greatest affinity scores were found for CDC2-like kinase 4 (CLK4, -8.4±0.0 kcal/mol), oligomeric death domain complex (CRADD/PIDD, -8.3±0.1 kcal/mol); progestogen receptor (PGR, -8.1±0.1 kcal/mol); and estradiol 17-beta-dehydrogenase 1 (17-beta-HSD 1, -8.1±0.1 kcal/mol). Proteins involved in endocrine disruption and breast cancer-related processes. Validation showed our protocol predicted the binding site for the 17-beta-HSD and PGR native ligands, in agreement with crystallographic data. Moreover, TCS interacts in the same binding site occupied by the 17-beta-HSD 1 native ligand, but in a different one from that used by progesterone in PGR. Protein-ligand binding involves hydrophobic and hydrogen bonding interactions. These results suggest TCS may target different proteins, probably altering pathways involved in pathophysiological disorders. Vice-Rectory for Research.


Over the last ten years there has been increasing interest in the identification and characterisation of tobacco smoke toxicants. We propose the use of a biologically relevant risk assessment framework incorporating both in vivo and in vitro data for the prioritisation of such toxicants. We have previously described the use of in vivo data in the generation of Margin of Exposure (MOE) values alongside Mode of Action (MOA) reviews. We have also proposed that individual toxicants are tested for activity in a battery of in vitro assays including in vitro micronucleus, Ames and mouse lymphoma assays.

MOE assessments are used as an initial tool to segregate tobacco smoke toxicants into high or low priority for risk reduction research. As recommended by EFSA, MOE values above 10,000 be considered a low priority for risk management actions. We have generated in vitro MOEs for a number of tobacco smoke toxicants in conjunction with MOA reviews and in vivo MOEs (where suitable data is available). Where the in vitro MOEs generated for individual toxicants do not provide a conclusive evaluation and are split across the critical value of 10,000 (e.g. NNK and arzonic) or where the available in vivo data is unsuitable for MOE generation (e.g. hydroquinone and catechol), the use of in vitro data can provide an alternative source of information. We present here MOE data for five different tobacco smoke toxicants:

Benz[a]pyrene: in vitro 42,469–3.0 x 10^7; in vivo 16,805–2.4 x 10^7.

NNK: in vitro 1.0 x 10^8–3.7 x 10^10; in vivo 338–3.7 x 10^10.

Asenin: in vitro 2.95 x 10^10; in vivo 13–4.9 x 10^9.

Catechol: in vitro 938–12,094; in vivo No data.

Hydroquinone: in vitro 1651–10,304; in vivo No data.
The incorporation of in vitro data into our suite of assessment methods allows us to generate additional evidence to support the 10 MAHs and improve the mechanistic understanding of individual toxicants. There is also the potential to incorporate such in vitro data into the future development of PBPK models for individual toxicants.


J. McKone, A. N. Van, E. Lachenaere and D. Johnson, University of California Berkeley, Berkeley, CA.

Orofacial clefts (OFCs) are caused by malformations in the closing of the lip or the soft palate. Although OFCs are considered to be one of the most common congenital birth defects, the underlying mechanisms of its etiology have yet to be clearly elucidated. This study sought to systematically determine how candidate genes previously identified in the literature interact, potentially implicating specific pathways for nonsyndromic OFC formation. OFC candidate genes were derived from genome-wide association studies (GWAS), the Comparative Toxicology Database, and Thomson Reuter's GeneGo. The compiled OFC genes were analyzed in conjunction with genes associated with the metabolic pathways of vitamin A and folic acid, factors shown to be implicated in OFC formation. Utilizing GeneGo and the candidate gene list, a gene-gene interaction network was constructed. Analysis of the network led to the identification of the TCF/LEF gene family, members of which regulate several downstream targets identified in the GWAS and/or are associated with vitamin A or folic acid metabolism. The TCF/LEF genes were shown in the network to be regulated by genes involved with the WNT signaling pathway, suggesting a mechanistic relationship between WNT signaling and orofacial cleft formation. Through computational analysis, this study proposes a potential gene-gene interaction mechanism for OFC formation via the WNT signaling pathway, which regulates several downstream targets identified in the GWAS and/or are associated with vitamin A or folic acid metabolism. The TCF/LEF genes were shown in the network to be regulated by genes involved with the WNT signaling pathway, suggesting a mechanistic relationship between WNT signaling and orofacial cleft formation.

122 Role of Beta-Methylamino-L-Alanine in GRK1-Mediated Amyotrophic Lateral Sclerosis Disease Pathway.

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Amyotrophic Lateral Sclerosis (ALS) is a debilitating neurodegenerative disease that affects about 30,000 people in the United States alone. Only a small fraction of cases are linked to genetics, making it likely that the overwhelming majority are induced by environmental factors. Therefore, the elucidation of a toxicity pathway is critical to understanding the vast majority of the ALS disease population. Amongst potential environmental exposures, neurotoxin beta-N-Methylamino-L-alanine (BMAA) has shown strong association with ALS in a small disease cluster in Guam. However, several proposed mechanisms are insufficient in explaining the entire disease pathway. In this study, we utilized computational predictions with GeneGo, STITCH, BLAST, PubChem, and KEGG to generate the most feasible pathway for BMAA toxicity. The results from PubChem structure clustering between a highly kainate-specific ligand and BMAA-beta-carbamate revealed a statistically significant 3D Tanimoto score, implicating a comparable receptor repertoire. BLAST sequence comparisons between GRK1-5 and GLUL, an ALS associated gene, showed significant overlap between the two nucleotide sequences, further suggesting that kainate receptors are crucial to the ALS disease pathway. Incorporating these computational studies with existing knowledge, there is compelling evidence suggesting a primary disease pathway in which BMAA-beta-carbamate binds to the kainate receptor GRK1 at the presynaptic motor neuron, leading to increased glutamate release and subsequent neurodegeneration.

123 Computational Analysis of a Potential Mechanistic Relationship between Depleted Uranium (DU) Exposure and Risk of Spina Bifida Cystica.

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Prolonged weapon use in war zones increases the environmental levels of potentially teratogenic metals which may have a direct relationship to increased incidences of birth defects. In Fallujah, Iraq since 2003, high levels of depleted uranium (DU) in the environment and in hair samples of residents suggests that constant exposure to DU may play a role in the alarming rates of congenital defects found in Iraqi children. One particular birth defect, Spina bifida, is a congenital defect of the spinal column with two major subcategories: spina bifida occulta and spina bifida cystica. Cystica is the most severe form of the disease but also the least prevalent. However, this has not been the case in Fallujah, Iraq where from 2003 cystica has been found to be the most prevalent form of spina bifida. The purpose of this study was to search for any commonalities between the genes and biological pathways associated with exposure to uranium and development of spina bifida cystica. Computational methods and tools utilized included the Comparative Toxicogenomics Database (CTD), PubChem, ToxNet, Pathway Commons, and unpublished exposure/defect data from Iraq. Out of the 292 genes reportedly affected by uranium exposure, 155 were also associated with spina bifida cystica. Biological pathway analysis revealed three genes, NT5E, PED, and TALDO1, associated with uranium and are also involved in upstream pathways of folic acid synthesis. Research has shown that a lack of folate is known to be related to spina bifida. This study provides a broad look at the genetic and molecular similarities between response to uranium exposure and spina bifida cystica and may offer guidance for future inventions. Further studies examining gene ontologies of populations exposed and not exposed to depleted uranium would offer a clearer picture of the relationship between uranium and spina bifida.

124 Computational Analysis of Catechins in Teas and Potential Relationship to Cardiovascular Health.

C. Lei, M. Nhan, C. Ha and D. Johnson, University of California Berkeley, Berkeley, CA.

Consumption of green tea has been reported to have beneficial effects on a variety of health-related issues. Of interest is the beneficial differences between green, oolong, and black teas, all produced from the leaves of Camilla senesis. Of the five main groups of bioactive compounds in the Camilla sinensis plant, catechins were found to be the most bioactive. Of the six main catechins in tea, epigallocatechin-3-gallate (EGCG), the most abundant and bioactive of the catechins was found to be the highest concentration in green tea as compared to oolong and black teas and subsequently was considered to be the most health-relevant component. The potential health benefits of tea depend on their level of fermentation, which is correlated to the amount of oxidation the catechins undergo during processing. EGCG is most abundant in green tea, which is not fermented, and highest in fresh unprocessed leaves from Camilla senesis. Green tea extract has also been reported to have increased biological activities. EGCG and prodrugs have also been shown to be potential epigenetic modulators. DNA methylation is catalyzed by DNA methyltransferase (DNMT) with S-adenosyl-methione (SAM) as the methyl donor. EGCG has been shown to reduce DNMT indirectly by reducing SAM and increasing S-adenosyl homocysteine (SAH) and homocysteine levels in the MCF-7 breast cancer cell line. In this study, primarily utilizing Genego Metadrug and MetaCore we report a proposed overlap between EGCG affected pathways and the atherosclerosis disease pathway. EGCG was found to competitively block hydroxy-3-methyl-glutaryl-CoA, and non-competitively inhibit squalene epoxidease, two rate limiting enzymes in the cholesterol biosynthesis pathway. In addition, previous animal studies have reported a decrease in LDL levels in rats given the three teas with the highest response coming from green tea treatment. This study provides an analysis of antioxidant compound levels found in tea and their relation to its effect on cardiovascular health, providing a possible explanation to the differences in beneficial effects found in different types of teas.

125 Computational Analysis of Environmental Factors Potentially Associated with Multiple Sclerosis Susceptibility.


Multiple Sclerosis (MS), an autoimmune disease of the CNS, may have associated environmental risk factors. A relationship between distance from the equator and MS risk reinforces the belief that UVB radiation and subsequent Vitamin D synthesis act protectively. The Human leukocyte antigen B chain, HLA-DRB1 is activated by Vitamin D and is a MOE to support the hypothesis that Vitamin D may offer protection from MS. HLA-DRB1*15:01, has been associated with MS susceptibility. Computational methods utilized to explore environmental factors beyond UVB radiation that might influence MS disease pathways included the Comparative Toxicogenomics Database (CTD), STITCH, GWAS data, GeneGo, and the EPA Toxic Release Inventory (TRI). Historical MS cases in the US were identified and toxic release data obtained through TRI Lead (Ph 5+), was found as a top contaminant among the MS clusters. Although lead has been proposed as a risk factor for MS, no previous published studies have clearly linked lead and MS. To identify and investigate genes,
126 Profiling the Activity of Environmental Chemicals in Causing Testicular Dysgenesis Syndrome Using the US EPA Toxicity Reference Database (ToxRefDB).

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Hyponadism, cryptorchidism, hypospadias, and testicular cancer are increasingly common male reproductive defects. Clinical and experimental evidence suggest that these defects are associated with testicular dysgenesis syndrome (TDS). We hypothesize an Adverse Outcome Pathway (AOP) framework for TDS starting with disruption of cell signaling and structural targets of the fetal testis during early-dysgenesis, followed by a series of key events that lead to male reproductive defects. Since understanding chemical-endpoint associations is an approach in building the AOP conceptual models, we mined 4209 guideline animal studies in EPA's ToxRefDB, which included 963 compounds tested in rats, mice, and rabbits. Testicular neoplasia was an outcome for 60 (6.2 %) chemicals. Sperm abnormalities were an outcome for 72 (9.8 %) chemicals tested in chronic and subchronic toxicity studies and 44 (5.7 %) chemicals tested in prenatal developmental and/or multigenerational fertility studies. Reductions in anogenital distance were recorded for 15 (3.5 %) chemicals tested in multigenerational studies, and 11 (1.5 %) chemicals tested caused hypospadias or cryptorchidism in development or multigenerational studies. The chemical classes showing the broadest activity across different TDS endpoints were pesticides (anilino, benzyol, primisulfuron-methyl, tetra-chlorobenzene, and vinclozolin) and phthalates (butyl benzyl phthalate, dibutyl phthalate, and diethylhexyl phthalate) with a lowest effect level on an of -300 mg/kg/day. The most potent TDS-actives (17 compounds) were determined as positive, and 11 or less of PENMC were calculated by ChemDraw® and Chem3D® for total 64 compounds using photochemical property including HLG. In this research, we established the hierarchical pattern of male reproductive defects in ToxRefDB which can be used as an AOP anchor for TDS. [This work does not reflect EPA policy].

127 Compound Toxicity Profiling and Prioritization Using Tox21 Phase I Quantitative High-Throughput Screening (qHTS) Cytotoxicity Data.

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Using in vitro data to prioritize compounds for in vivo toxicity testing is a goal of Tox21. Phase I profiled 1408 NTP compounds against 126 cell-based qHTS assays; those that measured cytotoxicity record the ability of a chemical to adversely perturb multiple cellular pathways in a concentration-response fashion. The curves from 39 cytotoxicity assays (9 human cell types, 4 rodent cell types, 13 sets of identical twin lymphoblastoid cell lines) were curated to filter noise and remove artifactual effects. To quantify the activity of each compound in these assays, we calculated a weighted version of Area Under the Curve (wAUC) intended to capture potency and efficacy simultaneously, as well as the conventional half-maximal activity concentration (AC50) value. The average wAUC or AC50 values across the 39 cytotoxicity assays were used to rank compounds and the rankings were compared, for 880 compounds, with available rat acute oral toxicity data. The compounds were categorized as toxic or non-toxic based on various “Globally Harmonized System of Classification and Labelling of Chemicals” (GHS) acute toxicity thresholds (50, 300, 500, 2000 mg/kg). The wAUC approach consistently prioritized more toxic compounds in the early stages of datasets at all thresholds. The best receiver operating characteristic (ROC) enrichment value at 1% and AUC value are 1.43 [95% CI=1.8-26.8] and 0.73 [95% CI=0.65-0.80], respectively, at a threshold of 50 mg/kg. Also, some toxic compounds (e.g., actinomycin D, chloric, daunomycin) had higher ranks based on wAUC due to their ill-fitted curves in some of the cell lines, resulting in missing AC50 values. We conclude that the wAUC provides an additional and useful metric to estimate the toxicity potential of compounds for Tox21 qHTS assays. Based on this metric, compounds screened in Tox21 can be prioritized for more extensive testing.

128 Predicting Cellular Dynamics and Key Events in Developmental Toxicity with a Multicellular Systems Model.


Computer simulation of cellular networks is one possible solution for modeling key events in developmental toxicology. We constructed a multicellular agent-based model (ABM) of early limb-bud development in CompCell3D (www.compueelld.org). The model simulates key cellular behaviors (mitosis, apoptosis, adhesion, migration, chemotaxis, shape, secretion), organizing centers (AER, ZPA) and signals (FGFs, SHH, BMPs, RA). It effectively emulates hindlimb-bud development during a 42h period in mouse (Thelier stages 16-19) and 160h in human (Carnegie stages 13-16). The ABM reflects biological variability across parallel simulations for spatio-temporal expression of biochemical gradients and cell behaviors, ultimately manifesting in trajectories of outgrowth. To evaluate the model as a tool for predictive toxicology, we selected 5-Fluorouracil (5FU) as a prototype. 5FU perturbed 13 of 650 ToxCast assays based on AC50s (or LECs) at or below 15 uM. 5FU effects observed in the assays were disruption of stem cell (mES) growth and differentiation, suppression of TGFβ1 signaling and mitochondrial density, p53-induction, mitotic arrest, reduced cell proliferation and increased cell death. Challenging the ABM with concentration-response data derived from mES cell number profiles, a dose-dependent wave of mitotic arrest and apoptosis, disrupting outgrowth. Varying the dose and time of exposure localized the primary key event to arrest of SHH-expressing cells and their geometric relationships to cells expressing GREM1, a BMP antagonist maintained by SHH signals. Different outcomes emerged when perturbation of the SHH/GREM1/BMP loop was switched between mitotic arrest and excessive apoptosis, indicating the importance of considering both cellular consequences together. These findings support the application of multi-cellular ABMs as tools to translate cellular dynamics into simulation of emergent (higher order) tissue effects for predictive toxicology. [This abstract does not necessarily reflect EPA policy.]

129 Chemical Structure-Based In Silico Phototoxicity Prediction: An Approach from a Combination of Photophysical Properties.


Some photophysical chemical properties are essential factor for prediction of phototoxicity, since phototoxicity is caused by photo-activation of the compounds. Highest Occupied Molecular Orbital – Lowest Unoccupied Molecular Orbital Gap (HLG) is a photophysical property of needful energy for photo-activation, and HLG was reported to be related with phototoxicity based on in vitro 3T3 Neutral Red Uptake assay (3T3 NRU assay). However there are few reports to predict of phototoxicity using photophysical property including HLG. In this research, we established the stepwise approach using Maximum-Conjugated Doublet-Electron-Number (PENMC) of the compounds in addition to HLG for in vitro phototoxicity prediction. HLG and PENMC were calculated by ChemDraw® and Chem3D® for total 64 compounds which were known the results of 3T3 NRU assay (32 positive and 32 negative). As step 1, we set the early cut line of PENMC as follows; the compounds that have over 8.0 of HLG were determined as negative, and the compounds that have less than 5.2 of HLG were determined as positive. On the other side, the compounds that have PENMC were calculated by ChemDraw® and Chem3D® for total 64 compounds which were known the results of 3T3 NRU assay (32 positive and 32 negative). As step 2, we set the early cut line of PENMC, respectively, at a threshold of 50 mg/kg. Also, some toxic compounds (e.g., actinomycin D, chloric, daunomycin) had higher ranks based on wAUC due to their ill-fitted curves in some of the cell lines, resulting in missing AC50 values. We conclude that the wAUC provides an additional and useful metric to estimate the toxicity potential of compounds for Tox21 qHTS assays. Based on this metric, compounds screened in Tox21 can be prioritized for more extensive testing.
was validated by the following results; sensitivity (84.4%), specificity (81.3%), positive rate (83.9%) and concordance (82.8%). We concluded that the stepwise approach with combination of HLG and PENW for prediction of phototoxicity is adaptable and useful for drug development as in silico screening, because it showed high sensitivity and negative rate from only chemical structure.

The current draft of the International Conference on Harmonisation (ICH) M7 guidance describes the use of in silico models to qualify genotoxic impurities during the drug safety evaluation and approval process. In order to attain the highest accuracy and improve the domain of applicability of the current Salmonella mutagenicity quantitative structure-activity prediction models, continuous updates to the training set must be made to accommodate new genotoxic findings. In this study, we first assessed our current Salmonella mutagenicity training set using fingerprints of known genotoxic structural alerts to determine the domain of applicability and performance characteristics of several commercial (QSAR) models. We then enhanced the previous version of our non-proprietary training database for Salmonella mutagenicity with data for 43 new compounds harvested from FDA approval packages and the published literature, to give a total of 3965 compounds. Of the 431 chemicals, 247 are drug molecules marketed between 1970 and 2011. Data gaps within the training set were identified and, using structural features derived from known toxicophores, 141 examples containing functional groups such as aromatic amino acids, hindered epoxides, propiolactones, amine halides, diazines, azo compounds, diazoniums, sulfates, aziridine chlorides, nitrites, hydrazines, nitriles, isocyanates, and sulfur mustard were added. Moreover, the new training set was expanded to include over 40 compounds containing previously unmodeled atoms such as boron, silicon, selenium, and tin. A hierarchical clustering analysis of the final training set showed representation of an additional 44 structural clusters over which the model can make a prediction.

The skin serves as a barrier against hazardous agents in the environment. However, the barrier is incomplete in that chemicals may penetrate the skin and cause toxicity. The absorbed dose depends not only on the absorption rate but also on the evaporation rate (assuming a fixed dose and skin area). The limited data available suggest that rates vary by several orders of magnitude between chemicals. However, there is a huge lack of data and new approaches are needed. One attractive possibility is to use quantitative structure-permeability relationship (QSPR) models. The aim of the present study was to examine different QSPR models addressing evaporation rate. We calculated evaporation rates for nine volatiles (methanol, n-propanol, acetone, methyl ethyl ketone, hexane, n-heptane, octane, benzene and toluene) at three air velocities and three air temperatures, using four semi-empirical models; McCready & Saghir, EPA, Mackay & Matsugu, and BAU. The predictions were compared with experimental data published by the EPA. None of the four models were able to predict the evaporation rate at all wind speeds. For comparison, we also developed linear solvation energy relationships (LSER) and partial least squares projections to latent structures (PLS) models. Seven solvents were used for calibration and two to test predictive performance. The LSER and PLS models showed good correlation (R2 0.95 and 0.92) and predictive performance (Q2 0.86 and 0.95) and seem more suitable than the semi-empirical models to predict evaporation rate. The semi-empirical models, EPA and Mackay & Matsugu at higher velocity (5.08 m/s) and lower temperature (280.35 K), showed a fair agreement between predictive and experimental values. As a result, evaporation rates can be adequately predicted from available physicochemical properties of the volatile organic compounds (VOCs). Reference 1. Braun, K.O. and Caplan, K.J. (1989) Evaporation rate of volatile liquids, final report, 2nd edition. EPA/744-R-92-001, NTIS PB92-232305, Springfield.

Cyclin Dependent Kinase Inhibitor 2A (CDKN2A) is a tumor suppressor protein in humans. It is capable of inducing cell cycle arrest in the G1 and G2 phases. p16INK4a is a major component of the RB pathway. p14ARF is part of an ARF-MDM2-TP53 system that exercises a negative control on hyper-proliferative signals originating from oncogenic stimuli. ARF binds to MDM2 and blocks its cytoplasmic transfer and thereby sequesters it in the nucleus. This hinders the MDM2 action, thereby blocking degradation of p53 and thus enhancing transcriptional activity.
and apoptosis. The purpose of this study was to identify additional regulatory targets of CDKN2A in lung epithelial cells, using Bayesian networks. Using the Robust Multi-Array Average procedure, a compendium of lung epithelial cell microarray data was generated based on Gene Expression Omnibus datasets GSE19027 and GSE994. The best-scoring regulatory networks, given the gene expression data, were then learned using the Bayesian Network Inference with Java Objects tool (BANJO) for static Bayesian Network inference. A set of known regulatory relationships involving CDKN2A was used as the initial network to focus the search space. Two proposers concluded in predicting IL1RN, IL6ST, IL1RAPL1, IGFAIS, and WNT10B as regulatory targets for CDKN2A. These genes are known to interact with a range of chemicals, including certain environmental toxicants. Furthermore, CDKN2A was predicted to be a direct regulator of RB1. These hypotheses warrant additional study as they lend valuable insights into the functions of CDKN2A in Chronic Obstructive Pulmonary Disease (COPD) and cancer. E2F3 was predicted, and validated in the literature, to be a regulator of MDM2 and TP53. Thus, Bayesian networks are a valuable tool for drawing signaling insights from gene expression data.

135 Irreversible Inhibition of Acetylcholinesterase by Soman.
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Acetylcholinesterase (AChE) is a key enzyme in the cholinergic nervous system that hydrolyses acetylcholine and terminates synaptic signals. Organophosphate compounds, such as nerve agents, can covalently inhibit AChE by phosphorylating the enzyme's catalytic serine residue. The phosphorylated adducts can either be reactive or be transformed to some limited extent by nucleophile compounds with oxime functional unit or undergo an aging reaction. Phosphonylation and subsequent aging leads to irreversible AChE inhibition, resulting in overstimulation of the nervous system. By employing ab initio QM/MM molecular dynamics simulations with umbrella sampling, we characterized the phosphorylation reaction mechanism between AChE and the nerve agent soman (GD), as well as the aging mechanism of GD phosphorylated AChE. The phosphorylation reaction between AChE and GD follows an associative nucleophilic substitution mechanism that is initiated when the nucleophile Ser200 attacks GD's phosphorus atom, with His440 acting as a general base. In the elimination step, Try121 of the catalytic gorge forms hydrogen bonds with the leaving fluoride atom prior to its dissociation from the active site. Once a stable covalent adduct is formed, the aging reaction begins with excision of the alkoxyl covalet bond connecting the bound GD's alky group to the phosphonate moiety. This cleavage is swiftly followed with a methyl group rearrangement of the alkyl group resulting in a stable tertiary carbenium, which is hydrated by an alcohol by a reactive water molecule facilitated by Try121. The characterized mechanisms and simulation results provide new detailed insights into this important process. Such mechanistic details are of significant interest for the development of novel strategies to reduce the toxic effects of GD poisoning by facilitating the search and design of novel compounds capable of slowing the aging of nerve-agent-inhibited AChEs as well as effective reactivators for the aged conjugates. Lastly, this work may also facilitate the design of aging-resistant pseudo-catalytic scavengers capable of sequestering nerve agents.

136 Computational Elucidation of Energetic Trends for DNA Intercalation.
The prediction of general energetic trends from intercalation of polyacrylamid compounds with DNA can provide insight into these pertinent interactions from a pharmacological and toxicological perspective. Compounds chosen for this study are nuclear base pair analogs (AT, GC, AA) and two novel intercalating agents (4-aza-tryptanthrin and coraline). Unfortunately, neither of these compounds have an established crystal structure when intercalated into DNA. The absence of this computational starting place gives rise to the need for a strong alternative computational method. Since dispersion forces have been shown to drive optimal orientation based on π-stacking, this method utilizes the electrostatic potential (ESP) maps of both interacting species. Presented herein is our improved method that uses more pertinent variables, overcomes the weakness of a center of mass method, and improves the efficiency with which computational studies can provide insight to experimental design. This approach relies primarily on visual intuition of the ESP alignments. The simplicity of this study leads to the decrease in the time cost and the consideration of dispersion forces increases confidence in our results. Our computational methods utilize SPARTAN and GAUSSIAN software, a Ground State Density Functional Theory (DFT), Local Spin-Density Approximation (LSDA) method, and a 6-31+G(dp) basis set. Initial results indicate a preferential intercalation of 4-aza-tryptanthrin with the GC base pair over an AT base pair, based on π-stacking. Identification of trends such as this can be valuable in designing future experimental studies targeting GC or AT rich regions associated with genetic disorders such as Myotonic Dystrophies, Fragile X, and Friedreich's Ataxia.

137 Efficient In Vivo Developmental and Neurotoxicity Screen of ToxCast Phase I and II Compounds in Zebrafish.
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The United States Environmental Protection Agency launched the Toxcast project to begin to predict the potential chemical toxicity of 1,078 compounds made up of pesticides, pharmaceuticals, “green” chemicals, chemicals in cosmetics and other consumer products. Early life stages are often sensitive to chemical insult, which make embryonic zebrafish an ideal platform to investigate the developmental and neurotoxicity of these compounds. We developed an efficient in vivo phenotypic screen using embryonic zebrafish to assess all 1,078 compounds. Using a wide range of concentrations (0.0864 to 64 μM, 10 fold serial dilution); all compounds were assessed for developmental toxicity beginning at 6 hours post fertilization (hpf). We kept exposed embryos completely in the dark until 24 hpf, and assessed photo-motor responses using the Photo-motor Response Assessment Tool (PRAT) that we developed. PRAT quantifies individual embryonic photo-motor response following two pulses of bright light. The initial pulse normally results in pronounced movement, and the second light pulse usually produces no activity. At 120 hpf, using Viewpoint Zebrabox, we assessed photo-induced larval locomotor activity. The locomotor activity is tracked for 25 minutes (10 in the light, 10 in the dark, and 5 in the light). Afterwards, each larva was assessed for changes in a suite of 20 morphological endpoints. We have successfully conducted the phenotypic screen on all 1,078 compounds, and a summary of the results will be discussed. Collectively, we have demonstrated the efficiency of the zebrafish model as a phenotypic screening platform to identify hazardous chemicals. This research is supported by NIEHS grants P30 ES00210 and R24ES019764.

138 In Silico and In Vitro Analyses of the Hormonal Activity of Hydroxylated Polychlorinated Biphenyl on Human Thyroid Receptor.
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Hydroxylated polychlorinated biphenyl (OH-PCBs) may disrupt thyroid hormone status because of their structural similarity to thyroid hormone. However, the molecular mechanisms of interactions with thyroid hormone receptors (TRs) are not fully understood. The integrated application of omics studies, bioinformatics, and computational modeling can provide to biological systems an enhanced understanding of the mechanisms underlying the toxicity of endocrine disruptor chemicals and support the study of disease etiology and prevention. In the present study, we examined the interactions between OH-PCBs and TRs to identify critical structural features and molecular properties of OH-PCBs related to their hormonal activity and to develop quantitative structure–activity relationship (QSAR) models for the thyroid hormone activity of OH-PCBs. Molecular descriptors were computed, selected, and used to characterize the ligand-receptor binding, and subsequently develop an in silico model. The in silico model had good robustness, predictive ability, and mechanism interpretability. Lipophilic distribution, hydrophobic and electrostatic interactions between OH-PCBs and TRs are important factors governing thyroid hormone activities. The OH-PCBs with higher ability to accept electrons, ortho position of the hydroxyl group, low dipole-dipole interactions tend to have weak binding with TRs and subsequently lower thyroid hormone activities. Hence, this in silico model can be used as a screening tool for further targeted toxicity testing and risk assessment, generate hypotheses about potential mechanistic pathways leading to adverse outcomes, and reduce time and cost of OH-PCBs testing.
A Systematic Analysis of ToxCast In Vitro Assays Associated with In Vitro Hepatic Outcomes.

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The U.S. EPA's ToxCastTM program uses hundreds of high-throughput in vitro assays to screen chemicals for bioactivity. The EPA Virtual Liver project combines ToxCastDB data with guideline rodent toxicology data to map hepatic adverse outcome pathways (AOPs). Here our objective is to reveal meaningful relationships between in vitro assays and in vivo liver outcomes. 289 ToxCast phase 1 chemicals were organized into 15 categories based on 272 histopathological lesions reported in guideline testing studies from ToxRefDB. Based on lesions reported at the study end, 242(83.7%) chemicals produced hypertrophy, 39(13.5%) chemicals resulted in hyperplasia, and neoplastic lesions were found for 79(29.4%). We compared the 150 chemicals with highest potency and the 150 lowest potency in the ToxCast assays and identified assays with at least 10 chemicals per category and the mean potency was significantly different (p<0.05). Out of the 973 assays, 28/147(19%) of the high-content imaging assays (APR) had associations with 7 liver injury categories. 6/83(7.2%) multiplexed transcription reporter assays (ATG) and 15/112(13.4%) protein complementation (OT) assays were also significant for liver injury. A number of the assays such as, cell loss, enzyme induction and enzyme inhibition, were generally significant across most categories. On the other hand, some assays were quite specific to certain categories. For instance, APR StressKinase_7hr_pos (stress kinase pathway activation) and ATG_CRE_CIS (transcription factor DNA binding activity) were significant for chemicals that showed regeneration vs ATG_VDRE_CIS (nuclear receptor transcription activity) for hypertrophy. The results suggest the utility of HTS assays for screening hepatotoxicity, and for linking biological events in the pathways to specific adverse outcomes. Such assays are not only mechanistically relevant but also aid in defining cell-based toxicity signatures for rapidly screening thousands of environmental chemicals. Abstract does not represent EPA policy.

Interpreting QSAR Toxicity Predictions in Hazard Assessments: An Acrylamide Case Study.


Toxicity predictions based on QSAR are used to screen large numbers of chemicals with little to no toxicity data into prioritized categories for further toxicity testing, development as “green” chemicals, or candidates for efficacious drug therapy. As databases incorporate more information on adverse pathways and potential targets, QSAR results may also contribute to hazard assessment for chemicals considered “data rich” (based on the availability of a full suite of traditional animal bioassays), but which are often “data poor” when it comes to characterizing the mode of action (MOA), severity progression, or potential adverse human clinical effects not monitored in animal bioassays. Presented here is an exercise to evaluate QSAR results within the context of extant hazard assessments on a data rich chemical, in this case, acrylamide (AA), and its less well studied active metabolite, glycidamide (GA). Results from four QSAR programs – two freeware programs (VEGA and OECD’s QSAR) and two proprietary programs (MetaCore™ and Discovery Studio/TopKat™) provided considerable new information in the areas mentioned above. Potential AA and GA induced adverse effects of clinical relevance to humans were identified as well as likely biological targets for AA, GA, and analogous chemicals. The results further supported the qualitative characterization of the MOAs, potential endpoints to consider in dose-response arrays, and identification of potentially important early biomarkers to aid future bioassays or human study design. Conversely, extant AA and GA ADME data and PBPK models helped assess the likelihood of AA or GA reaching biological targets predicted by the QSAR programs. This exercise demonstrated the value of integrating QSAR results into traditional hazard assessment for improved qualitative information. The next step is to integrate high throughput screening/content (HTS/HTC) assay results into the assessment to further quantitate dose-response for a wider array of effects. [The views expressed are those of the authors and do not necessarily reflect the views or policies of the U.S. EPA]

Health-Related Effects Reported by Electronic Cigarette Users in Online Forums.

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Electronic cigarettes (e-cigarettes) are battery-operated devices that deliver aerosolized nicotine to users without burning tobacco. Because little data exists on their health effects, we explored the symptoms that e-cigarette use has on humans by analyzing online user posts from three e-cigarette forums with “health and safety concerns.” Basic information (location, age, and gender) and health (symptoms and doctor diagnosed signs) information were collected. A total of 405 symptoms (78 positive, 326 negative and 1 neutral) were reported in three forums. Most data analysis was performed on Electronic Cigarette Forum (ECF) posts. A total of 12 systems/anatomical regions were affected in e-cigarette users. Systems most often affected include: mouth and throat, respiratory, neurological, sensory, and digestive. The majority of negative health effects occurred in the respiratory system. We further consolidated reported symptoms into categories to determine which anatomical regions/physiological processes were most affected for each system. For consolidated data, symptoms were most frequently reported for: bronchi/lungs (e.g., wheezing, loss of breath, difficulty breathing, throat), gastrointestinal (headaches), intestine/digestion, and sight. To analyze interactions between systems, interactions were created with Cytoscape software. Interactions were most frequently seen between circulatory/neurological; respiratory/mouth and throat; respiratory/chest, and digestive/neurological systems. Increased blood pressure was the most frequently reported sign diagnosed by physicians treating e-cigarette users. While some positive health effects were reported, a significant proportion of the data showed a correlation with e-cigarette use and onset of adverse health effects. This study is the first to compile and quantitatively assess health data associated with e-cigarette use from online forums.

The Relationship Between T-Box Transcription Factors CAMK2B and PITX2 in the Expression of Collagen Protein COL4A3 in COPD.


The T-Box transcription factors are known have recently been highlighted for their expression role in epithelial cells in Chronic Obstructive Pulmonary Disease (COPD), which genes, including COL4A3, TBX2, TBX3, TBX5, PML, CFLAR, GULP1, CASPIO, PAX3, BOK, and PITX2 are connected in transcriptional regulatory networks of lung epithelial cells. The type IV, alpha 3, collagen gene (COL4A3) involves into extracellular matrix construction as the major structural component of basement membrane. COL4A3 is known to have suppressed expression in COPD3. The A polymorphism of the COL4A3 is also indicated of associated with the risk of developing COPD. COL4A3's C terminus binds to autoantibodies at basement membranes in Goodpasture syndrome and is phosphorylated by calcium/calmodulin-dependent protein kinase II beta (CAMK2B), which is activated by the promyelocytic leukemia protein (PML). The paired-like homeodomain 2 (PITX2) regulates the expression of N-cadherin. N-cadherin changes the adhesion of extracellular matrix (ECM) by interacting with collagen proteins. PITX2 gene is regulated by different kinase pathways, such as MAPK and Akt. However, the relationship between these factors is still not well studied. Thus, we have come up with the question that if is PITX2 gene also regulated by CAMK2B. We conducted the study in focus on exploring the relationship between PITX2 and CAMK2B genes by applying Bayesian Network Structural Learning (BNSL). Version 2.2 of Bayesian Network Inference with Java Objects (Banjo) was employed in the study. The result was used to compare the known interactions between COL4A3, CAMK2B, PITX2 and PML. The study has shown the possible novel phosphorylation of PITX2 by CAMK2B.

Validation of a Systems Toxicology Based Adverse Outcome Pathway Prediction with Functional Outcome: Effect of Exposure to 2, 4-Dinitrotoluene on Energy Metabolism and Exercise Endurance.

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2,4-dinitrotoluene (2,4DNT), commonly used in industrial and explosive manufac- turing processes, is known to contaminate manufacturing areas and munitions manufacturing facilities leading to its listing on US EPA's Contaminant Candidate List. Previous transcriptomic and lipidomic studies identified energy metabolism as a potential target of DNT toxicity. The impact of such perturbations of energy metabolism on exercise endurance is largely unknown. We hypothesized that organism-level impacts of 2,4DNT dosing were the result of energy metabolism deficits, especially lipid metabolism, from involving interference with PPARalpha signaling and its downstream pathways. To validate this adverse outcome pathway, we linked molecular changes caused by 2,4DNT exposure to effects...
on whole animals. PPARα (-/-) and wild-type (WT) mice exposed to a sublethal 2,4-DNT dose (134 mg/kg/day for 14 days) or vehicle were given an exercise challenge (a forced swim) 1 day after the last dose to determine how 2,4-DNT and/or PPARα impairment affected overall performance. Observations were collected at multiple levels of biological organization including genes involved in fatty acid and glucose metabolism, PPAR activation and response, biochemistry (serum triglycerides and glucose), and swimming endurance. Decreased swim times were observed with DNT in WT and PPARα (-/-) mice, but DNT effect was significantly less in knock-down mice indicating that knock down of PPARα expression partially rescued mice from DNT-induced energy metabolism deficits. Our results support the proposed hypothesis by demonstrating that 2,4-DNT's impact on energy metabolism, especially lipid metabolism, occurs via perturbation of PPARα signaling resulting in reduced exercise endurance at the individual level. (Support: US Army Corps of Engineers)

**144 Modelability of ToxCast Phase I Datasets.**
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One of the problems in Quantitative Structure-Activity Relationships (QSAR) analysis is to establish, whether it is possible to build a predictive model for a given dataset. For some datasets, all attempts to build a predictive model using different sets of descriptors and QSAR/QSTR methodologies fail raising a question, whether it is possible to evaluate the dataset modelability prior to modeling. We have devised several modelability criteria such as dataset diversity, new activity cliff indices, correct classification rate (CCR) for similarity search models (sCCR), CCR=0.5 (sensitivity=specificity), etc. These criteria were applied to 40 binary datasets, for which QSAR models were built using Dragon 5.5 descriptors and/or ToxCast in vitro assays treated as biological descriptors, and KNn, Random Forest and SVM methods. The best modelability criterion was found to be the sCCR, which had the correlation coefficient of 0.73 with the QSAR/QSTR model CCR. We consider a model predictive, if its CCR as well as both sensitivity and specificity are at least 0.70. We found that to satisfy this condition, sCCR should be at least 0.68. sCCR values were obtained for ToxCast datasets with 24 ToxRefDB in vivo assays as end points, which had at least 30 toxic compounds among 212 compounds of the curated ToxCast dataset. None of the sCCR for these datasets was as high as 0.60 except for the rat cholinesterase inhibition assay, for which it was 0.85, and sensitivity and specificity were 0.83 and 0.88, respectively. We conclude that with the latter exception, ToxCast Phase I datasets with ToxRefDB in vivo assays as end points do not appear to be modelable using QSAR approaches. This conclusion agrees with the recent empirical observations of Thomas et al. (Toxicol Sci. 2012, 128:398-417).

**145 Celestrol Decreases Specificity Proteins (Sp) and Fibroblast Growth Factor Receptor-3 (FGFR3) in Bladder Cancer Cells.**
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Bladder cancer is the ninth most common cancer worldwide and ranks 13th as a cause of cancer deaths. MVAC (methotrexate, vinblastine, Adriamycin and cis-platin) chemotherapy has been extensively used for treatment of advanced bladder cancer and is accompanied by toxic side effects and, thus, it is important to develop less toxic alternate chemotherapeutic therapies and dietary management strategies for prevention of this disease. Celestrol (CSL) is a naturally occurring triterpenoid acid that exhibits anticancer activity, and in K7 and 253JB-V bladder cells, 0.5 to 2.5 μM CSL induced apoptosis, inhibited growth, colony formation and migration, and CSL decreased bladder tumor growth in vivo. CSL also decreased expression of specificity protein (Sp) transcription factors Sp1, Sp3 and Sp4, and several Sp-regulated genes/proteins including vascular endothelial growth factor, survivin and cyclin D1. CSL also decreased fibroblast growth factor receptor-3 (FGFR3), a potential drug target for bladder cancer therapy. Results of mRNA interference and knockdown of Sp proteins show that FGFR3 is a Sp-regulated gene downregulated by CSL. The mechanism of Sp downregulation by CSL was cell context-dependent due to activation of proteosome-dependent (KU7) and -independent (253JB-V) pathways. In 253JB-V cells, CSL induced reactive oxygen species (ROS) and inhibitors of ROS such as glutathione blocked CSL-induced growth inhibition and repression of Sp1, Sp3 and Sp4. This response was due to induction of the Sp repressors ZBTB4/ZBTB10 through downregulation of miR-27a and miR-20a/17-5p, respectively, which regulate expression of these transcriptional repressors. Thus, the anticancer activity of CSL in 253JB-V cells is due to induction of ROS and ROS-mediated induction of Sp repressors ZBTB4/ZBTB10 through downregulation of miR-27a and miR-20a/17-5p and this is emerging as a characteristic pathway observed for many ROS inducers in cancer cells.

**146 Raloxifene Potentiates the Cytotoxicity-Induced by RL91, a Second Generation Curcumin Analog, in PC3 Prostate Cancer Cells.**
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The survival rate for men with hormone refractory prostate cancer has not changed significantly in the last 30 years. Thus there is a need for new drug treatments for this aggressive cancer. Our lab has had success with second generation curcumin derivatives as novel therapies for aggressive breast cancer. In this study we examined the combination of raloxifene and 2,6-bis-(pyridin-4-ylmethylene)-cyclohexaneone (RL91), a potent 2nd generation curcumin derivative as a novel treatment for hormone refractory prostate cancer (HRPC). The combination treatment showed highly potent cytotoxicity toward PC3 prostate cancer cells compared to individual treatments. Specifically, EC50 values of 2 μM and 10 μM were produced by RL91 and raloxifene, respectively. Moreover, this combination decreased cell number by 85% compared to control after 96 h of treatment, as determined by the sulfonrodamine B assay. Raloxifene is known to modulate the activity of estrogen receptor alpha (ERα) and beta (ERβ). The activation of these receptors as well as the epidermal growth factor receptor (EGFR), is crucial for the progression of HRPC. To determine how raloxifene potentiates the cytotoxic effect of RL91, the localization of these receptors was examined by fluorescence microscopy. The results showed that ERα, ERβ, the androgen receptor (AR) and EGFR were expressed in PC3 cells. However, raloxifene treatment (10 μM for 48 h) promoted EGFR internalization in the cytoplasm. A similar effect was also seen for ERα where raloxifene promoted a translocation from the nucleus to the cytoplasm. However, no change was observed for either ERα or the AR. These results suggest that raloxifene-mediated changes in the localization of ERβ and the EGFR provide a mechanism by which raloxifene enhances the cytotoxicity of RL91 toward PC3 cells. This novel mechanism should be explored further in order to develop new therapies for HRPC.

**147 Evaluation of Wild Yam (Dioscorea Villosa) Root Extract As a Potential Epigenetic Agent in Breast Cancer Cells.**
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Aberant epigenetic alterations in the genome, is believed to be a potential cause of some forms of cancer. Due to their reversibility, epigenetic modifications are considered potentially useful in drug development approaches (epi-drugs). The current available synthetic epi-drugs are non-specific and induce adverse effects. Natural products might offer advantages and find utility for cancer treatment. The present study was designed to evaluate the efficacy of wild yam root extract as a potential demethylating agent using two breast cancer cell lines, MCF-7 (Estrogen receptor positive, ER+) and MDA-MB-231 (ER negative, ER-), and a gene, GATA-3, a potential marker of breast cancer development. Moreover, GATA-3 expression is methylation-specific, being higher in ER+ cells with promoter hypomethylation and insignificant in ER- with promoter hypermethylation. In this study, cells, approximately at 70% confluency, were treated with wild yam root extract (0-50 μg/ml) for 72h and then used for viability, mRNA, and methylation analyses. It was observed that wild yam significantly reduced viability of both cell lines and enhanced the mRNA contents of DNMTs (DNMT1, 3A, and 3B) and GATA-3 in a dose-dependent manner. Global DNA demethylation, as measured by 5′-deoxytetraycytidine (mC) and 5-hydroxymethylcytosine (hmC), showed that mC was increased only in MCF-7 cells, whereas hmC level was reduced in both cell lines. Since hmC is generated from mC by ten-eleven translocation (TET) enzymes, the present data suggest that enhanced expression of GATA-3 and DNMT enzyme mRNAs followed by a reduction in mC in MCF-7 and MDA-MB-231 cells are the mechanism of suppression of TET enzyme functions in the epigenome by wild yam root extract. This plant with a long history of traditional use should be further explored with regard to its potential as an epigenetic agent in breast cancer therapy.
Commensal bacterial community shifts in the pathogenic colonic environment and chronic colonization of mucosa-associated Escherichia coli (MAEC) has been linked to colonic tumourigenesis. Enteropathogenic Escherichia coli (EPEC) is one of the most commonly identified MAEC in colorectal cancer patients. The aim of this study is to address the contribution of MAEC colonization to human carcinogenesis. EPEC infection of cancer cell caused alterations in affect locomotion-related behaviors. Of cancer cell including detachment, migration, cytokinetic rearrangement, dissemination and survival via induction of macrophage inhibitory cytokine 1 (MIC-1). Mechanically, MIC-1 induced RhoA GTPase which mediated survival of the detached cancer cells. In terms of signaling pathway, MIC-1 triggered TGF-beta-activated kinase 1 (TAK-1), which enhanced expression of RhoA GTPase. In conclusion, mucosal EPEC enhanced MIC-1 gene expression in the human intestinal cancer cells, which was associated with enhanced tumor cell resistance to anoikis and subsequent survival via enhanced TAK-1 and RhoA GTPase. This work was supported by the Basic Science Research Program through the National Research Foundation of Korea, funded by Ministry of Education, Science, and Technology Grant 2012R1A1A2005837.

**Ring-Substituted Analogs of 3, 3'-Diindolylmethane (DLM)**

**Induce Apoptosis and Necrosis in Androgen-Dependent and -Independent Prostate Cancer Cells.**

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We have recently reported that novel ring-substituted analogs of 3,3'-diindolylmethane (ring-DIMs), exhibit anti-androgenic and anti-proliferative activities in androgen-dependent prostate cancer cells. We hypothesized that the anti-proliferative effects of ring-DIMs may be due to their ability to induce cell death. Ring-DIMs inhibited androgen-stimulated LNCaP cell proliferation and induced apoptosis and necrosis in LNCaP and PC-3 prostate cancer cells with 2-4 fold greater potency than DIC. The cytotoxicity of the most potent ring-DIM, 4,4'-dibromo-DIM, but not the other ring-DIMs, was decreased by caspase-3 inhibition. The 4,4'-dibromo-DIM was found to be specifically taken up by the extracellular matrix, whereas the ring-DIMs were intracellularly distributed in both cell lines. Ring-DIMs were more potent inhibitors of cell growth and survival in LNCaP and PC-3 cells than DIM and the differential structure-dependent cell death mechanisms indicate ring-DIMs have clinical potential as chemopreventive and chemotherapy agents in prostate cancer, regardless of hormone-dependency.

**Dichloroacetate (DCA) Increases Radiation Sensitivity of A549 and H1299 Lung Cancer Cells.**

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Dichloroacetate (DCA) is a synthetic small molecule inhibitor of pyruvate dehydrogenase kinase used to treat metabolic diseases with low toxicity in human patients for decades. Recent reports suggest that DCA is cytoxic to several cancer cell types including non-small-cell lung cancers (NSCLC). The cytotoxicity to cancer cells appears to be a result of reversal of the Warburg effect through partial restoration of mitochondrial glucose oxidation away from cytoplasmatic aerobic glycolysis. With the increase in glucose oxidation, ROS production is augmented leading to the opening of the mitochondrial transition pore to release cytochrome c and apoptosis inducing-factor (AIF) into the cytoplasm. The consequence is mitochondria membrane depolarization and induction of cell death in a caspase-dependent or -independent manner. Given the potential therapeutic translation of DCA to human cancer treatment, we investigated the efficacy of DCA to sensitize NSCLC to X-ray induced killing in vitro after both single and fractionated X-ray exposure. Treatment with DCA decreased plating efficiency (PE) and enhanced radiation-induced cell killing by dose modifying factors of 1.5 in A549 and 1.4 in H1299. The decrease in PE was not due to induction of apoptosis or necrosis. X-ray fractionation showed that DCA inhibited split-dose recovery/repair by 3.5 fold in A549 and 1.5 fold in H1299. Flow cytometry analysis with propidium iodide indicated significant cell cycle arrest at G1S in A549 and at G2/M in H1299. The data suggest that DCA enhances X-ray-induced NSCLC cell killing through inhibition of DSBR and/or alteration of cell cycle distribution.

**Mango Polyphenolics Reduce Inflammation in Intestinal Colitis—Potential Involvement of the miR-126/Pi3K/Akt/mTOR Pathway In Vitro and In Vivo.**

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Mango polyphenolics including gallic acid, mangiferin and galloolaminoids, have shown antioxidant, anti-inflammatory and anticarcinogenic properties in several studies. However, anti-inflammatory mechanisms relevant to the prevention of colon cancer have not been well investigated. This study investigates the potential role of the miRNA-126/Pi3K/Akt/mTOR signaling pathway in the anti-inflammatory effects of mango polyphenolics in human CDC-18Co colon-myofibroblastic cells and on DSS-induced colitis in rats. Animals were administered control juice (15.7 g sugar and 0.05 g citric acid/100 ml) or mango juice (total phenolic content of 475.80 mg/L GAE), and exposed three cycles of 3% DSS. The mRNA and protein levels were measured by RT-PCR, western blot analysis and immunohistochemistry. In vitro mango extract and gallic acid suppressed the expressions of inflammatory mediators such as NF-kB (p65) and IL-1β and reduced the expressions of AKT and HIF1α involved in AKT/mTOR pathway at mRNA and protein level in a dose dependent manner. Correlatively, miRNA-126, which negatively regulates the Akt/mTOR pathway, was induced by mango extract and gallic acid. In the rat colitis model, mango juice intake suppressed cell proliferation as measured by Ki-67 staining, and resulted in protection against DSS-induced colon inflammation during chronic colitis compared to control juice. The juice significantly attenuated the expressions of pro-inflammatory cytokines such as TNF-α, IL-1β, IL-6, and IL-10 at protein and mRNA level. Moreover, the phosphorylation of AKT and mTOR were suppressed, and the expression of Pi3K was reduced while miRNA-126 that has a target site in the mRNA of Pi3K was upregulated by the juice.
These results suggest that mango polyphenols attenuated inflammatory response by down-regulating the expression of pro-inflammatory cytokines and up-regulating the expression of anti-inflammatory cytokines. This finding is in line with previous studies that have reported the anti-inflammatory effects of mango polyphenols in various models of inflammation

153 Loss of the Tumor Suppressor Protein PTEN Contributes to Increased Nrf2 Signaling.

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PTEN is a master regulator of cyto-protective genes involved in the maintenance of cellular redox balance. In recent studies, the P38/Akt pathway is reported to positively regulate the PTEN pathway. The protein phosphatase PTEN counteracts the action of PI3K by removing the phosphate from PIP3. This led us to hypothesize that loss or mutation of PTEN, an event that occurs frequently in cancers, might increase Nrf2 activity that promotes the survival and proliferation of cancer cells. In our study we investigated the role of PTEN in prostate and breast cancer, commonly reported to have either loss of heterozygosity or mutations in PTEN. Using the PTEN-null human prostate cancer cell line PC3, we found that expression of wild-type but not mutant PTEN decreased the basal and antioxidant induced expression of Nrf2 target genes. More common clinically, patients have loss of a single allele of PTEN. Therefore, we also investigated the Nrf2 pathway in cells heterozygous for the PTEN gene. Using a murine prostate tumor cell line with a spontaneous loss of a single allele in PTEN (PTEN-P8, +/-) and an isogenic cell line that had deletion of the second allele (PTEN-CaP8, +/-) we found that complete loss of PTEN resulted in higher expression of the Nrf2 target genes. In order to fully address the role of PTEN haploinsufficiency that could mimic the cellular progression from benign to malignant we also looked at Nrf2 activity in the immortalized breast epithelial cell line MCF-10a that had both alleles intact (+/+), deletion of a single allele (+/-), or deletion of both alleles (-/-) of PTEN. As expected, we found that both the basal and inducible expression of the prototypical Nrf2 target gene NQO1 inversely correlated with PTEN status. Together this suggests that the loss of PTEN contributes to increased Nrf2 transcriptional activity, which likely provides cells with an altered proteome that favors oncogenesis.

154 Formaldehyde-Induced Replication Stress Causes Activation of ATR Kinase Leading to p53-Mediated Apoptosis and Senescence in Human Lung Cells.

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Formaldehyde (FA) is a recognized human carcinogen with documented inhalation exposures in many occupational groups. Here we examined stress signaling pathways and cell fate decisions triggered by FA in human lung cells (normal lung fibroblasts, H460 lung epithelial cells and primary bronchial epithelial cells). We found that FA induced a rapid activation of the DNA damage-responsive ATR kinase that preferentially targeted the p53 transcription factor at low and moderate damage levels. Activation of p53 was evidenced by its strong Ser-15 phosphorylation, protein stabilization and upregulation of its target genes, such as MDM2 and CDKN1A (p21). Knockdown experiments with siRNA confirmed the p53 dependence of the gene expression responses. FA also caused a depletion of the p53 inhibitor MDM4 via its enhanced proteolysis. The use of biochemical markers of individual cell cycle phases and replication status manipulations showed that activation of ATR-p53 signaling by FA occurred exclusively in the S phase. The phase may play a major role in FA-induced apoptosis in lung epithelial cells, which was associated with upregulation of the proapoptotic gene BAX (BCL2L1). The presence of p53 was also required for permanent growth arrest (senescence) in FA-treated lung fibroblasts. Overall, our results indicate that replication stress and ATR-activated p53 are responsible for cytotoxic responses in FA-exposed human lung cells. Acknowledgements. This work was supported by grant ES020689 from NIEHS.

155 Down-Regulation of Telomerase Activity and Shortening of Telomere Length by Polychlorinated Biphenyls (PCBs) in HL-60 Cells.

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We reported that PCBs, environmental persistent organic pollutants and probable human carcinogens, can down-regulate telomerase activity in vitro in immortal human keratinocytes (HaCaT). The most efficacious conger was the dioxin-like PCB126, a potent arylhydrocarbon receptor (AhR) agonist. To analyze whether this effect occurs in vivo and is tissue specific, we examined the mode of action (MOA) we exposed another cell type, human promyelocytic leukemia (HL-60) cells, to PCB126 and also to PCB153, not an AhR activating congener. Both compounds reduced telomerase activity, visible after 6 days of exposure, and telomere length, to about 50% within 30 days of exposure, PCB126 more so than PCB153. This reduction in telomerase activity and telomere length was seen in both cell lines, but only HaCaT also showed a strong increase in cytochrome P450 1A1 mRNA and activity, the hallmark of AhR activation. This suggests that AhR activation may be one, but not the only mechanism for this effect of PCBs. HL-60 can differentiate which is accompanied by a reduction in telomerase activity. However, the continued proliferation of the PCB-exposed cells makes this mechanism less likely. Telomeric repeat binding factors (TRF1, TRF2) are involved in the stabilization of telomers. Up-regulation of TRF1/2 was seen in PCB126-exposed HaCaT, pointing to a possible mechanism. Experiments with both cell lines, representing different target tissues of PCB toxicity, are under way to elucidate the MOA of PCBs on telomeres and the possible significance of this effect on precursor cells of the hematopoietic pathway. (Supported by NIEHS P42 ES013661)

156 Chronic Exposure to Particulate Hexavalent Chromium Disrupts Shugoshin1 Localization in Human Lung Cells.

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Chromosomal instability (CIN) is a hallmark of cancer and can be caused by spindle assembly checkpoint disruption or chromosome missegregation during mitosis. Hexavalent chromium (Cr(VI)) is a well-known human lung carcinogen, and has shown to induce numerical CIN, however its mechanisms for inducing aneuploidy remain unknown. In this study we are investigating whether Cr(VI) affects a key centromeric cohesion protein, Shugoshin 1 (Sgo1). Sgo1 maintains and protects centromeric cohesion in G2 and continues to maintain proper sister-chromatid cohesion during mitosis. This protection mechanism prevents sister-chromatid separation prematurely during mitosis. Disruption of Sgo1 localization has been shown to lead to chromosome missegregation. We have found that chronic exposure to particulate Cr(VI) disrupts the localization of Sgo1 in G2 cells. Specifically, after a 24 h exposure to 0.1, 0.2, or 0.3 µg/cm2 lead chromate we did not observe any changes in the percent of G2 cells with Sgo1 localization at the kinetochores. However, a 120 h exposure to the same concentrations showed a concentration-dependent decrease in the percent of G2 cells with Sgo1 localization at the kinetochores. Specifically 0.1, 0.2 or 0.3 µg/cm2 disrupted localization in 36, 88, and 90% of G2 cells, respectively. Our findings suggest that particulate Cr(VI)-induced CIN is mediated through disrupting Sgo1 localization at the kinetochores during G2 cells, thus leading to chromosome missegregation and ultimately aneuploidy. This work was supported by NIEHS grant ES016893 (J.W.).

157 Ultraviolet B-Irradiated L-Tryptophan Induces Human UDP-Glucuronosyltransferase 1A1 and 1A8 Expressed in the Human Skin.

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Benz[a]pyrene is a widespread environmental contaminant and its active metabolite, benz[a]pyrene-7,8-dihydrodiol-9,10-epoxide, is associated with the development of ultraviolet (UV) B-induced skin cancer. While UDP-glucuronosyltransferase (UGT) is involved in the detoxification of benz[a]pyrene, the protective role of UGT in the defensive mechanism against UVB-induced skin cancer has not been elucidated yet. Here, we investigated the effects of UVB irradiation on the expression of human UGT1A enzymes as well as on cytochrome P450 (CYP) 1A1 in HaCaT cells. Multiple UGT1A isosforms such as UGT1A1, UGT1A3, UGT1A4, and UGT1A8 were expressed in human skin and HaCaT cells. When HaCaT cells were treated with 6-formylindolo[3,2-b]carbazole (FICZ), which is one of the tryptophan derivatives formed by UVB, UGT1A1 and UGT1A8 along with CYP1A1 were significantly induced. While UVB-irradiated tryptophan also induced those enzymes, the formation of FICZ was not detected in the irradiated tryptophan solution, indicating that tryptophan derivatives other than FICZ formed by UVB might have the potential to activate the aryl hydrocarbon receptor. While UVB induces CYP1A1, increasing the bioformation of benz[a]pyrene-7,8-dihydrodiol-9,10-epoxide from benz[a]pyrene, it also induces UGTs, accelerating the detoxification of benz[a]pyrene. Specific induction of skin UGT1A1 and UGT1A8 without inducing CYP1A1 might protect individuals from developing skin cancer.

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The eukaryotic translational initiation factor (eIF4E) is an essential component of the cellular translational machinery and is responsible for binding ribosomes to the cap structure of mRNAs. The phosphorylated form (serine-209) of eIF4E plays a critical role in cancer cell growth and transformation. Treatment of colon cancer cells with a synthetic cannabimimetic WIN 55,212-2 (WIN) inhibited cancer cell growth, induced apoptosis and downregulated specificity protein (Sp) transcription factors and Sp-regulated gene products associated with cancer cell growth (EGFR and Cyclin D1), angiogenesis (VEGF and VEGFR) and survival (survivin and bcl-2). The anticancer activity of WIN is accompanied by induction of multiple phosphorylations and some of the effects of WIN are blocked by the phosphatase inhibitor sodium orthovanadate (SOV). Treatment of SW480 cells with 7.5 μM WIN alone also decreased levels of eIF4E and co-treatment with 0.35 mM SOV blocked WIN-induced downregulation of eIF4E and knockdown of Sp proteins confirmed that eIF4E was an Sp-regulated gene. Protein phosphatase 2A (PP2A) catalyzes dephosphorylation of both eIF4E and Mnk-1, an upstream kinase that phosphorylates eIF4E. Treatment of SW480 cells with 7.5 μM WIN and knockdown of PP2A by RNA interference blocked downregulation of eIF4E, Sp proteins and some Sp-dependent genes. Several anticancer agents inhibit Sp transcription factors by inducing zinc finger binding protein ZBTB10 and by suppressing microRNA-27a (miR27a). Treatment of SW480 cells with 7.5 μM WIN induced ZBTB10 protein and decreased miR27a expression and knockdown of PP2A reversed these responses demonstrating that WIN-induced downregulation of Sp and eIF4E was due to PP2A-mediated disruption of miR27a/ZBTB10 axis.

Recent studies have demonstrated that non-coding RNAs (ncRNAs) are differentially expressed and play an important role in gene regulation and influence normal and cancer cell phenotypes. About 3000 IncRNAs have been identified and some of these act as scaffolds regulating molecular (protein, RNA and DNA) interactions required for various signaling networks and this is accomplished, in part, by association with chromatin-modifying complexes. HOXA transcript at the distal tip (HOTTIP) is a 3,764-bp IncRNA transcribed from 5’ tip of HOXA gene cluster and is expressed in anatomically distal cells such as hand and foot fibroblasts. Previous studies in our laboratory showed that HOTAIR is a negative prognostic marker in pancreatic cancer and this is due, in part, to interaction of HOTAIR with Polycomb Repressive Complex 2. In contrast, HOTTIP was reported to interact with MLL complexes by specifically binding to WDR5 adapter protein leading to broad loss of H3K4me3 across HOXA locus. In this study, we investigated the functional role of HOTTIP in Panc1 and L3.6P.1c pancreatic cancer cell lines. Knockdown of HOTTIP by RNA interference studies (iHOTTIP) resulted in ~50% reduction in cell survival within 72 hr after transfection and iHOTTIP also decreased pancreatic cancer cell migration which was determined in a Boyden chamber assay. iHOTTIP enhanced Annexin V staining and PARP cleavage associated with induction of apoptosis in both Panc1 and L3.6P.1c cells. Knockdown of HOTTIP also decreased expression of cyclin D1, VEGF and survivin which play an important role in cell proliferation, angiogenesis and survival. An in vivo study was performed using L3.6P.1c cells transfected with iHOTTIP in a mouse xenograft model and after 15 days of knockdown there was a significant reduction in tumor volumes and weights (670%). RNAseq analysis of HOTTIP knockdown in Panc1 cells resulted in significant (p<1.5 fold) changes in expression of ~1467 genes and analysis of data suggests that HOTTIP mediated gene regulation has a critical role in pancreatic cancer progression.

Anoikis is an anchorage independent cell death. Resistance to anoikis is one of the key features of the metastatic cells. In the current study, we have shown the role of STAT-3 in anoikis resistance in various melanoma and pancreatic cancer cells. Marked anoikis was induced as such in the cells that were grown in anchorage-independent conditions. The melanoma and pancreatic cancer cells that resisted anoikis were observed to have higher rate of migration as compared to the cells that were exposed to anchorage dependent growth, as observed in cell migration and invasion assays. These anoikis-resistant cells also had significantly higher expression and phosphorylation of STAT-3 at Tyr 705, than the cells that were attached to the basement membrane. Treatment of these cells with IL-6, a cytokine which phosphorylates STAT-3, prevented the induction of anoikis. STAT-3 inhibitors AG490 and pipilartine induced anoikis in a concentration-dependent manner, whereas IL-6 blocked anoikis. Over-expression of STAT-3 by transfection, not only increased the anoikis resistance but also protected cancer cells from pipilartine-induced anoikis, confirming the role of STAT-3 in anoikis resistance. On the other hand, silencing STAT-3 decreased the potential of cancer cells to resist anoikis. Furthermore, STAT-3 (+) cancer cells were more sensitive to anoikis as well as to the effect of piplartine as compared to STAT-3 (-) cells. The STAT-3 (+) cells also had enhanced migration potential as compared to STAT-3 (-) cells. In summary, our results establish STAT-3 as a critical player that renders anoikis resistance to the cancer cells and enhance their metastasis potential. The role of STAT-3 in anoikis resistance in breast and ovarian cancer is currently under investigation. The outcome of this study has great clinical implications as it will serve as a platform to devise rational therapeutic approaches to treat metastatic cancers. [Supported in part by R01 grants CA106953 and CA129038 (to S.K.S) awarded by the National Cancer Institute].

Indole-derived compounds including diindolylmethane (DIM), indole-3-carbinol, tryptophan photoproducts and metabolites have been identified as agonists of the aryl hydrocarbon receptor (AHR). Indole and tryptophan metabolites tryptamine (TA) and indole 3-actetate (IAA) are produced by gut microflora and there is evidence that AHR ligands modulate gut inflammatory pathways. We therefore investigated the AHR agonist/antagonist activities of indole, TA and IAA using Ah-responsive breast and colon cancer cells as in vitro models. In MDA-MB-468 breast cancer cells (TA and indole 3-actetate) and MDA-MB-231 breast cancer cells (IAA and IAA and IAA), the concentrations of IAA and IAA and IAA increased CYP1A1 mRNA and protein with an EC50 < 500 μM. IAA and indole also induced CYP1A1 mRNA (only IAA induced CYP1A1 protein) however < 30 and < 10% of the maximal induction responses were observed at 1000 μM concentrations of IAA and indole respectively. The effects of these compounds as inhibitors of 2,3,7,8-tetrachlorodibenz-p-dioxin
(TCDD) induced responses were also investigated in cells treated with the indole compound alone or in combination with TCDD. Indole (500 and 1000 μM) significantly inhibited TCDD-induced CYP1A1 mRNA and protein levels and similar results were observed for 500 μM TA however the former compound was a significantly more potent AHR antagonist. In contrast IAA did not exhibit AHR antagonist activity. These results were somewhat variable among different cell lines however, it was evident that the major gut microbiome product, indole, was an AHR antagonist and this may impact AHR-dependent gut inflammatory pathways.

**163 Tolfenamic Acid Inhibits Colon Cancer Cell and Tumor Growth and Downregulates Specificity Protein (Sp) Transcription Factors.**

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Introduction: Tolfenamic acid (TA) is a nonsteroidal anti-inflammatory drug (NSAID) and is a potential chemotherapeutic agent for treatment of colon cancer; however, the mechanism of action of TA is unknown and was investigated in this study.

Methods: Inhibition of colon cancer cell growth and induction of apoptosis by TA was investigated using cell counting and Annexin V staining and modulation of specificity protein (Sp) transcription factors Sp1, Sp3, Sp4 and Sp4-regulated gene products was determined by western blot analysis of whole cell lysates. Mechanisms of TA-induced Sp downregulation were investigated using specific pathway inhibitors and the in vivo antitumor activity of TA was examined in athymic nude mice xenografted using RKO cells as xenografts.

Results: TA induced apoptosis and decreased colon cancer cell growth and this was accompanied by caspase-dependent proteolysis of Sp1, Sp3 and Sp4 and decreased expression of Sp-regulated genes including bcl-2, survivin, VEGF, VEGFR1, cyclin D1 and c-MET. TA also inhibited colon tumor growth and decreased Sp1, Sp3, Sp4 and Sp4-dependent gene product expression in tumours.

Conclusion: TA-induced repression of Sp transcription factors and Sp-regulated genes play a role in the cancer chemotherapeutic effects of TA. Since TA acts as an anticancer agent in several tumor types, results of this study suggest for the first time that TA is a potential chemotherapeutic agent for treatment of colon cancer. Clinical applications of TA alone or in combination treatment of colon cancer are enhanced since this agent is relatively non-toxic and has previously been used as a non-steroidal anti-inflammatory drug. The prior use of TA, as an NSAID will also facilitate approval of the drug for application as a cancer therapeutic agent.

**164 Ni2+-Induced Chromosome Aberrations/Gene Amplification/Gene Silencing Alter Cytoskeleton, Ca2+ Distribution, and Global Gene Expression, Causing Morphol/Neoplasm. Transformation of 10T1/2 Mouse Embryo Cells.**

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Ni refinery workers inhaling Ni sulfide ore dusts/smoking cigarettes contracted lung/nasal cancers. Inhaled Ni3S2/green NiO induced lung cancer in rats. Ni3S2/green-black NiOs induced chromosomal aberrations/morph-neoplasms transformation (Tx) in 10T1/2 mouse cells. Ni/MCA-Tx cell lines showed a) ect-2 gene amplification/higher ect-2 mRNA/protein, b) no DRIP80 c) no β-centaurin-2 mRNA. We hypothesized Ni2+) 1) amplified ect-2 gene, causing higher levels of microtubules (MTs); 2) silenced β-centaurin-2 gene, causing higher levels of microfilaments (MFs); and 3) silenced DRIP gene, altering Ca2+ distribution/Tx 10T1/2 cells. We tested these hypotheses by staining cells with fluorophores (Fluo) to decorate MTs; then examined cells by confocal microscopy. In non-Tx 10T1/2 cells, MFs/MTs were arranged in long fibers. In Ni/G green Ni-Ox-Tx cell lines, MFs/MTs were over-expressed, aggregated in areas, absent/other areas, changing cell shapes. Low density non-Tx cells had high nuclear/low cytoplasm Ca2+ concentration (State I); high density near-confluent cells had low nu./high cyto. Ca2+ (State II). Ni/MCA-Tx cell lines were largely in State II. We conclude Ni2+ ions 1) amplified ect-2 silenced β-centaurin-2 genes, causing over-expression of MTs/MFs, altering cell shapes, changing global gene expression; 2) silenced DRIP80 gene, altering Ca2+ distributions in Tx cells; and 3) induced mutations/methylations in 15 genes, causing differential expression of 130 genes, contributing to induction/alteration of chromosome aberrations/morph-neoplasms in Ni2+/MCA-Tx cell lines.

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**165 Benzoquinone-Induced Topoisomerase Modifications: Linking Benzene Myelotoxicity and Leukemia.**

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Protein adduction by reactive electrophiles can induce structural and functional changes that contribute to toxicity and disease progression. Such electrophiles are often products of xenobiotic metabolism or generated endogenously via oxidative stress and lipid peroxidation. Relevant to this phenomenon are the redox-active and electrophilic metabolites of benzene, which are believed to contribute to its myelotoxic effects. When the benzene metabolites hydroquinone (HQ) and phenol (PHE) are administered to rats, HQ oxidizes to 1,4-benzoquinone (BQ) and in the presence of GSH gives rise to multi-GSH conjugates detectible in bone marrow. These HQ-GSH conjugates retain the ability to adduct proteins and to redox cycle. Here we report that bone marrow malondialdehyde levels in PHE/HQ treated rats are significantly elevated, indicative of lipid peroxidation and the consequent generation of other reactive electrophilic aldehydes, such as 4-hydroxy-2-nonenal (4HNE). Indeed, proteomics profiling revealed bone marrow proteins targeted by benzene metabolites and 4HNE, including 14-3-3 protein delta, protein disulfide isomerase a3, peroxisiredoxin 2, and calreticulin. Addition of topoisomerase II α (topo IIα) has been implicated in benzene-induced leukemia, and cancer chemotherapy topto IIα inhibitors are a leading cause of therapy-induced leukemia. We next reacted purified topo IIα (6 units) with BQ (0.5 μM) or 4HNE (1.3 μM). A marked reduction in the ability of topo IIα to decantate the kDNA substrate was observed. Proteomic analysis of 4HNE-reacted topo IIα revealed multiple amino acid sites of addition, including K953 and K1480 adducts. Addition of these lysine residues could impair topo IIα-DNA binding, or inhibit topo IIα’s APT-dependent formation and annealing of DNA strand breaks. The consequences of BQ- and 4HNE-induced functional alterations in topo IIα are currently under investigation.

**166 Overexpression of CRM1 in Normal Human Lung Epithelial Cells Changes Cellular Morphology and Cytotoxic Responses to Tobacco-Specific Carcinogen NNK.**

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Chromosome region maintenance 1 (CRM1), the major nuclear export receptor with a broad substrate range, is not only required for transport of many RNAs and proteins but also involved in various modulations within the cell such as mitosis, cell arrest, and apoptosis. Our recently published study showed that CRM1 played critical roles in response to tobacco-specific carcinogen, 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanone (NNK), in BEAS-2B cells (a normal human lung epithelial cell line). The objective of the present study was to further examine the significance of CRM1 in lung cancer development using BEAS-2B cells stably overexpressing CRM1 (BEAS-2BCRM1+ cells). The overexpression of CRM1 in BEAS-2BCRM1+ cells was confirmed by real-time PCR and western blot. As compared to BEAS-2B cells, BEAS-2BCRM1+ cells were prone to form colonies. Soft-agar assay further demonstrated increased colony formation and larger colony size in BEAS-2BCRM1+ cells in comparison with BEAS-2B cells. In addition, the cytotoxic effects in response to NNK was measured by 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium (MTS) assay in both cell lines. Cells were treated with 0.500 μM NNK for 24, 48, and 72 h. The inhibitory effects were dose and time dependent in both cells (p<0.05). However, BEAS-2BCRM1+ cells showed different sensitivity to NNK as compared to BEAS-2B cells. Taken together, our results indicate that CRM1 overexpression changes cellular morphology and cytotoxic response to tobacco carcinogen in human lung epithelial cells. The potential molecular mechanisms involving these changes are being evaluated to better understand the critical role of CRM1 in chemical carcinogenesis after NNK exposure.

**167 ARNT Isorforms Mediate Opposing Effects on NF-κB Signaling.**

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We have previously shown that the arylhydrocarbon receptor nuclear translocator (ARNT) regulates the chromatin binding activity of the RbB NF-κB subunit. ARNT is a transcription factor that is integral in the regulation of xenobiotic and hypoxic responses but our studies suggest that ARNT also participates in NF-κB
signaling. ARNT is expressed as two isoforms, isoform 1 and 3, but whether these isoforms differentially regulate NF-kB activity is unclear. Isoform 1 is identical to isoform 3 except for an additional 15 amino acids near the N-terminus. The extra amino acids in isoform 1 provide a phosphorylation site, which has been shown to inhibit its DNA binding. Interestingly, we have observed in lymphoid malignancies that ARNT isoform 1 is expressed at much higher levels than isoform 3 as compared to normal lymphocytes where the ARNT isoforms are expressed at equal levels. We find that ARNT isoform 1 potentiates while ARNT isoform 3 abrogates NF-kB activity. In light of our previous ARNT-ReB model, these opposing effects of the ARNT isoforms in NF-kB signaling appear to hinge on their ability to associate with RelB as ARNT isoform 3 binds much more strongly than isoform 1. Lastly, we found that the co-expression of a GFP-tagged p100 in combination with the ARNT isoforms promoted p100 nuclear translocation, possibly through a RelB bridge, resulting in diminished transactivation of NF-kB responsive genes. Thus, our current working model is one in which lymphoid malignancies shift toward the production of ARNT isoform 1, which in turn enhances NF-kB activity, as a contributing factor to their growth and survival.

168 Co-Exposure to Arsenic and Estrogen Leads to Enhanced Transformation of Normal Human Prostate Epithelial Cells.

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Exposure to both arsenic and estrogen are known risk factors for prostate cancer, however the carcinogenic effect of their co-exposure is not known. Therefore, the objective of this study was to evaluate the transformation potential and its mechanism in human prostate epithelial cells by co-exposure to these two chemicals. To achieve this objective, the human prostate epithelial cells, RWPE-1 were treated for 6 months with sodium-meta arsenite and 17β-estradiol, both alone and in combination, at concentrations of 100 pp/mL and 100 ng/mL. Cell counts and MTT assay was performed to determine the effects on growth, and soft agar assay was performed to evaluate the cell transformation by exposure to arsenic and estrogen. Potential role of estrogen receptors and aromatase in mediating the cellular response to these chemicals was evaluated by measuring their expression at transcript level. The result of this study revealed that the growth and transformation of RWPE-1 cells were significantly greater in arsenic and estrogen co-exposed cells as compared to the individual cells exposed to these two chemicals. The data of quantitative real time PCR revealed that expression of estrogen receptor beta was significantly increased whereas aromatase was significantly decreased in arsenic and estrogen co-exposed cells. These findings together with our recently published data on aberrant expression of epigenetic regulatory genes, such as, DNMTs, HDACs and MBDS in arsenic and estrogen co-exposed cells suggest that co-exposure to these two chemicals enhances carcinogenicity through epigenetic mechanisms.

169 Effects of Polycyclic Aromatic Hydrocarbons with Estrogen Receptors α and β in Breast Cancer Cells.

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Polycyclic aromatic hydrocarbons (PAHs) are widespread environmental pollutants found in cigarette smoke, contaminated soil, vehicle exhaust, among others, known to cause adverse health effects including carcinogenesis and endocrine disruption. The metabolites of PAHs have been implicated in endocrine disruption. Some PAHs are known to have estrogenic effects by interacting with estrogen receptors (ERs). Estrogens have diverse roles including regulation of mammary gland development and morphogenesis, and maturation of uterus and ovaries. These functions are mediated by two subtypes of estrogen receptors, ERα and ERβ, which function in different forms (i.e. ERαβ homodimer and ERαβ heterodimer) ERβ is known to be proliferative while ERβ is known to be antiproliferative in the mammary gland. The estrogenic effects of PAHs have been found to depend on their effects on ERs. Recently our lab reported that the monohydroxylated metabolites of naphthalene, phenanthrene and pyrene showed differential effects with ERα and ERβ. This study is essential to ensure effective therapy.

170 Alternative Splicing of ATG5 in DU145 Human Prostate Cancer Cells Inactivates Autophagy and Promotes Xenograft Tumor Growth.

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Autophagy is a highly conserved pathway that targets cytoplasmic cargo to the lysosome for degradation. Excess or aberrant organelles, large protein aggregates and non-selective portions of the cytosol can be targeted by the autophagosome and delivered to the lysosome in order to combat cell death or maintain cell survival in response to stress. In this study, we show that DU145 human prostate cancer cells alternatively splice Autophagy-related protein 5 (ATG5), an essential protein for autophagosome formation. These novel ATG5 splice forms are unable to conjugate normally to ATG12 and instead are rapidly ubiquitinated and turned over by the proteasome. The absence of ATG5-ATG12 conjugate in DU145 cells prevents autophagosome formation and autophagic degradation. Stable expression of full-length ATG5 using lentiviral infection rescued both ATG5-ATG12 conjugation and autophagy. Autophagy is currently thought to be tumor suppressive by eliminating potentially genotoxic protein aggregates and damaged organelles. However, the exact role it plays in tumor progression and survival is currently under investigation. To address this question, we performed subcutaneous injections of autophagy-deficient DU145 cells into immunodefficient mice and monitored tumor formation and growth. In initial experiments, autophagy deficient tumors had a significantly longer latency period, yet grew at a faster rate than autophagy competent tumors. This suggests that autophagy can promote initial tumor establishment, while suppressing later tumor growth. Currently there are numerous clinical trials investigating the therapeutic potential of drugs that inhibit autophagy. Since it’s not yet clear how these opposing aspects affect cancer metastasis, further study is essential to ensure effective therapy.

171 The Aryl Hydrocarbon Receptor Regulation during Epithelial-to-Mesenchymal Transition.

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The aryl hydrocarbon receptor (AhR) is a ligand-activated transcription factor that plays a role as a mediator of the xenobiotic signaling pathways. Recently, the AhR emerged as a major player in breast carcinogenesis. We have shown previously that ectopic overexpression of AhR in immortalized normal human mammary epithelial cells (HMEC) resulted in the development of malignant phenotypes, most notably the epimorphic transformation of HMEC. However, it is not rigid but is in a dynamic state; with cells moving back and forth between epithelial and mesenchymal states. We asked the question whether this flux of phenotypes is due to changes in AhR levels or activity. A clone of HMEC overexpressing AhR, with either epithelial (E) or fibroblastic (F) phenotypes was compared to an empty vector (EV) control cells. We employed Western blotting, immunocytofluorescence (ICF) staining and RT-PCR techniques to assess the AhR expression as well as the epithelial and mesenchymal markers, E-Cadherin and vimentin, respectively. Our results showed that although at the mRNA levels AhR expression was similar, the AhR protein was higher in E- than in F-type cells. The AhR protein levels were always higher when cells were grown at higher density. Nuclear localization as assessed by ICF and CYP1A1 expression as assessed by RT-real time PCR, were used as surrogates for AhR activation. Our analysis showed that AhR activation was much higher in F-type cells and is more pronounced at higher cell density. In contrast, a 3h treatment with TCDD, a potent AhR agonist resulted in a steeper induction of CYP1A1 in E-type than the F-type cells, which is further enhanced at higher density. The observed difference in AhR protein levels and activity may be due to a differential stability of the protein associated with the two morphological forms, which is under investigation.

172 Expression and Role of ALDH1B1 in Pancreatic Cancer.

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Recent studies show that ALDH activity selectively defines an enhanced tumor-initiating cell population in human pancreatic adenocarcinoma. The specific ALDH isozyme(s) contributing to the high ALDH activity in these cells are unknown. We have recently shown that the mitochondrial aldehyde dehydrogenase 1B1
ALDH1B1 is a potential biomarker for colon cancer. The aim of the current study was to understand the biology and the role of ALDH1B1 in pancreatic adenocarcinoma. In normal pancreas, ALDH1B1 is abundantly expressed in glandular cells, but sparsely in the ducts (ALDH1B1 immunopositivity = 16.7 ± 1.7). In pancreatic ductal carcinoma, we found a rather high ALDH1B1 expression in ductal cancerous tissues (ALDH1B1 immunopositivity = 197.2 ± 29.4). Our data were also confirmed by analyzing human pancreatic adenocarcinoma tissue microarrays. Furthermore, ALDH1B1 appears to be contributing to ALDH activity in some, but not all, human pancreatic cancer exfoliated samples. The variation of ALDH1B1 expression was also observed in 16 human pancreatic cancer cell lines. Two high and two low ALDH1B1 expressing cell lines were subjected to siRNA ALDH1B1 knockdown and cell proliferation was evaluated. High ALDH1B1 cell lines showed a 35% reduction in cell growth whereas proliferation was not affected in low ALDH1B1 cell lines. Treatment of high ALDH1B1 cells with gemcitabine resulted in greater ALDH1B1 mRNA levels and enzyme activity. In contrast low ALDH1B1 knockdown and cell proliferation was evaluated. High ALDH1B1 cell lines were subjected to siRNA (ALDH1B1) treatment with a luciferase reporter fused to a HIF Response Element (HRE) to determine down-regulation of the prometastatic gene CXCR4, previously been accompanied down-regulation of the prometastatic gene CXCR4, previously been reported as an AHR ligand-inducible gene. Omeprazole and related benzimidazole analogs also exhibited anti-invasion activities in vitro suggesting that this AHR-pharmaceutical class may have clinical importance for treating advanced basal-type breast cancer that metastasizes to other tissues. (Supported by NIH-R01-CA-136571).

### Gene Expression Changes in Human Endometrial and Mammary Cells Exposed to Tamoxifen

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Tamoxifen (TAM), a selective estrogen receptor modulator used for adjuvant therapy and/or prevention of breast cancer, also increases the risk of endometrial and myometrial cancer. We hypothesized that comparison of gene expression patterns in cultured normal breast and endometrial cells may elucidate TAM-induced mechanisms. Gene expression studies in normal human mammary epithelial cells (NHMECs) exposed to 10 μM TAM for 48 hr, using NIH DNA-oligonucleotide microarrays (Schild et al., 2005), demonstrated up-regulation of genes active in the interferon and immune-response pathways. Here we used the same NHMECs and TAM exposure conditions, and evaluated gene expression by Human Gene 1.0 ST Affymetrix expression array. Results using the Affymetrix array confirmed TAM-induced up-regulation of interferon signaling and complement pathways. Genes significantly up-regulated included some reported before (IFIT2, IFIT1, IFIT3, IFIT4L, IFITM1 and OAS3), and new genes involved in the same pathway (IFH4, IF35, and IFIH1), as well as genes in the complement system (C1S, C1R and SERPING1). To contrast breast with endometrium, we used human endometrial stromal cells (HESC, cells), also exposed to 10 μM TAM for 48 hr. The primary genes up-regulated included some involved with biosynthesis of steroids (DHCR7, FDPS and MVLD), and SREBF2, a sterol transcription factor. In addition, there was up-regulation of PARG, a gene involved in cell proliferation and cancer, and down-regulation of CC1, a gene with anti-tumor activity. Expression changes of the most highly altered genes have been confirmed by qRT-PCR, and the microarray data are being subjected to extensive pathway analysis. However, these preliminary data show induction of different gene expression patterns in normal human mammary and endometrial cells exposed to TAM, confirming that immune-response pathways are induced in the breast, and showing that proliferative and proinflammatory pathways are induced in the endometrium.

### Revealing the Role of Cancer Testes Antigens in HIF Signaling

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The hypoxia –inducible factor-1 (HIF-1), is a transcription factor that responds to changes in oxygen homeostasis. Furthermore, the HIF pathway has been observed to contribute to tumor aggressiveness. The overexpression of HIF proteins has been observed to increase invasive and metastatic behavior in tumor cells. Identifying the mechanism that regulates HIF is essential to developing therapeutic strategies to inhibit its function. Here, we asked whether any members of the cancer-testes (CT)-antigen (CTA) family support HIF signaling. We combined an siRNA mediated loss of function approach with a luciferase reporter fused to a HIF Response Element (HRE) to determine consequences on HIF signaling following individual depletion of 120 CTA. This then revealed a subset of CTA-antigens that positively support HIF induced transcription. This regulation appears to be at the level of the HIF1a protein as depletion of a subset of these CTAs, MAGEA3/6 and IF2BP3 reduced induction of HIF1α protein. These findings suggest that CTAs may support stress signaling, particularly under hypoxic conditions and further demonstrate that CTAs may be playing functional roles in supporting tumor cell survival.
Glycidol fatty acid esters (GEs), trace contaminants in edible oils which are possibly formed during refining processes, have recently been detected in vegetable fat-containing products, including infant formulas. The level of GEs was ten times higher in enzymatically processed dacylglycerol-rich oil than that in the regular cooking oil containing triacylglycerols as major components. Although there is no toxicological data available yet on the GEs, the primary toxicological concern is based on the potential release of genotoxic carcinogen, glycidol from the parent esters. In the present study, to detect the modifying effects of GEs on the mammary gland, one of the carcinogenic target organs of glycidol, we pretreated 7-week-old SD rats with N-methyl-N-nitrosourea (50 mg/kg i.p.) and then administered glycidol (800 ppm) or GEs (3600 ppm, glycidol oleate (GO) or glycidol linoleate (GL)) in the drinking water for 26 weeks. The dose levels being selected on the basis of carcinogenic dose levels in rat carcinogenicity study of glycidol (37.5 and 75 mg/kg/day) and on the equal moles of the esters. In body weights, significant decrease was noted in the glycidol group compared to control group from week 2 through experimental period due to obvious decrease of water consumption. The calculated glycidol intake was 43 mg/kg per day and on the assumption that all treated GEs would be completely metabolized to glycidol, intake of glycidol in GO and GL groups was 93 and 74 mg/kg per day, respectively. The multiplicity and volume of histopathologically diagnosed mammary tumors, in particular poorly differentiated carcinomas were significantly increased in the glycidol group as compared with the control. In the GO group, the multiplicity and volume of mammary tumors showed a slight tendency to increase, but no change was noted in the GL group. These results provide evidence of a mammary tumor promoting activity of glycidol, but not GEs in the present model.

**Spontaneous Thymoma Observed in Carcinogenicity Study of Wistar Han Rats.**

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Wistar Han rats are an appropriate model for toxicity and carcinogenicity studies in rodents and spontaneous thymoma is sometimes recorded in carcinogenicity studies of this strain of rats. The incidence of thymoma in historical control data of Harlan Laboratories is as follows: benign thymoma 0.64% for males, 2.39% for females; malignant thymoma 0.50% for males, 0.63% for females. The incidence of benign thymoma is higher in females with a range of 0% to 17.02% in these data. Histologically, these tumors commonly appear as solitary lesions with expansive growth, consisting of a mixture of thymic epithelial cells and lymphocytes with medullary differentiation. It was not always clear and requires careful consideration to distinguish between hyperplastic lesions and benign thymoma, and also between benign and malignant thymoma for many cases. In this report, we introduce typical hyperplastic lesions, thymoma and also the rarer epithelial cell type thymoma observed in Wistar Han rats.

**Mechanisms of Acetylenegenol Nanoparticles on Melanoma Development: In Vitro and In Vivo Assays.**

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Eugenol displays antiproliferative and pro-apoptotic activities in different types of cancer cells, although effects of Acetylenegenol (AC) and Acetylenegenol Nanoparticles (NCAG) have not been elucidated. Here the role of NCAG on in vivo melanoma model and their actions on in vitro melanoma and endothelial cell cultures were investigated. Mutine melanoma cells (B16F10, 8X105/100mL) were s.c. injected in the dorsal region of C57/Bl6 mice. Animals were i.p. or p.o. treated with Saline, AC, NCAG (50 mg/kg/day) or with their respective controls during 7 days. In vitro human endothelial cells (HUVEC) and melanoma cells (5x10^5/2mL) were incubated with RPMI, DMSO, NC, AC or NCAG. Cell viability, nitric oxide (NO), clonogenic survival and cell adherence were monitored. Only NCAG (100μM and 300μM), treatment reduced the endothelial and melanoma cell viabilities, but not AC and NCAG (60μM) treatments reduced the clonogenic survival only in melanoma cells and AC (30 and 60μM) treatment reduced clonogenic survival in both endothelial and melanoma cells. AC and NCAG (60μM) treatments inhibited endothelial and melanoma cells adherence. Both NC and NCAG (10, 30 and 60μM), but not AC, treatments increases the NO production by endothelial and melanoma cells. I.p. injection of NC or NCAG reduced NO production, nevertheless they caused loss of weight, due to lower food intake and reduced the number of platelets. These in vivo toxic effects may be caused by accumulation of NC or NCAG in the peritoneum. Do administration of NC or NCAG reduced melanoma growth more than AC, and NC, AC or NCAG treatments decreased the number of circulating leukocytes. Together, data obtained show that i.p route is not feasible to NC treatments, and the efficiency of NCs on tumor cell growth detected by p.o. may be due to their higher activity on clonogenic survival and adherence of melanoma cells.

**Natural Compounds As Chemopreventive Agents for the Inhibition of Protein Targets Involved in Cancer Induced by UV Radiation.**

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Skin cancer is one of the most common worldwide, with an increasing incidence in recent years. This disease is mainly caused by excessive exposure to solar ultraviolet (UV) radiation. Currently, there is a high demand for natural chemopreventive compounds that may work on the biochemical mechanisms involved in the development of the disease. In this study, in silico molecular protein-ligand docking was performed with Autodock Vina to assess the interaction of 44 natural bioactive compounds with protein kinases (ERK1/2, PK, p38/2YIX, and JNK2/2DDT) and cyclooxygenase-2 (3LN1), widely recognized protein targets in the signaling cascades of skin tumor formation induced by UV radiation. The results showed these compounds presented theoretical binding affinity scores of similar magnitude to those recorded for known inhibitors of these proteins. The best binding affinity values were further scored for binding activity compared to ERK and JNK (10.3±0.1 kcal/mol), and to cyclooxygenase-2 (9.7±0.1 kcal/mol). In the case of p38, the greatest affinity was found for epigallocatechin-3-gallate (-9.2±0.0 kcal/mol). The affinities obtained for the inhibitors of ITO, 2YIX, 2ZDT and 3LN1 were -9.4±0.0 (FR-180204), -7.0±0.0 (CE-159167), -7.3±0.0 (C46), and -10.3±0.0 kcal/mol (celecoxib), respectively. These theoretical results are good indicators that natural bioactive compounds may work as potential chemopreventive agents against skin cancer induced by UV exposure, probably by a mechanism involving their direct binding on key protein targets associated with the disease. Vice-Rectory for Research. UniCartagena. 2011-2012. Colciencias-UniCartagena, Colombia: Grants 11074592166 (2009) and 110751929058 (2010).

**Genetic Polymorphism of Human Microsomal Epoxide Hydrolase As a Determinant of Polyaromatic Hydrocarbon Metabolism and Toxicity.**

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Microsomal epoxide hydrolase (mEH, EPHX1) is a key catalytic determinant in the formation of diol-epoxide metabolites of certain polycyclic aromatic hydrocarbons (PAHs), noted as potent and ultimate mutagenic moieties. Epidemiological associations between EPHX1 genetic status and the incidence of certain diseases, including lung cancer, have been reported, yet the mechanistic bases for these associations remain unclear. Further, PAH diol-epoxides have never been evaluated as substrates toward the fjord region dibenzo[a,l]pyrene-11,12-diol-13,14-epoxides. Results remain unclear. Among the mEH variants, Y113/H139 wild type allele displayed highest capacity abilities to metabolize bay region and fjord region PAH diol-epoxide intermediates. Among the mEH variants, Y113/H139 variant exhibited lowest affinity for the bioactivation of the fjord region dibenzo[a,l]pyrene-11,12-diol-13,14-epoxides. Results
from Comet assays and in situ DNA damage assays conducted in COS1 cells trans-

cfected with the mEH variants and treated with the corresponding epoxides gen-
erally corroborated the enzymatic activity data. Overall, these findings demonstrate

marked substrate selectivity among the mEH variants with respect to PAH epoxide 

metabolism and provide mechanistic support for published epidemiology data sug-
gerusting that the H113 mEH allele is associated with a reduced risk of lung cancer.

182 Cotinine Levels and Gene Polymorphisms in Asthmatic Children Exposed to Tobacco Smoke in Northern Mexico.


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Tobacco smoke (TS) represents a serious health threat to consumers and passively 
exposed individuals. However, passive exposure receives much less attention than 

smokers. In addition, more than 700 million children worldwide are passively ex-

posed to TS, meaning an increased risk to develop tobacco related diseases later in 

life; among them children with respiratory impairments, like asthmatics, are a par-

ticularly vulnerable group since the lung is the main route of TS entrance. Asthma 
is a multifactorial disease and the environment quality is a relevant etiology factor. We 

investigated children exposure to tobacco smoke in an asthmatic population (n=100 

individuals; 6.98 ± 0.85 years), and non-asthmatic children (n=100, 7.1 ± 

0.82 years) living in Región Lagunera, Northern Mexico. There was a statistically 
significant difference in urinary cotinine between the exposed and non-exposed 
groups. Urinary cotinine levels showed a small positive correlation between chil-
dren and smoking adults (r=0.1205 and p= 0.244). Although, cotinine levels were 

higher in asthmatic children compared to non-asthmatic, such difference was not 

statistically significant. In addition, no association between cotinine levels and 

asthma severity was observed in this study. As for the allelic frequencies of GSTT1 

genotype, no significant differences were observed between the exposed and non-exposed 
groups. GSTT1 null and CYP2A6*2 (1799 T-A) polymorphisms, were significantly different 

comparing TS exposed healthy and non-asthmatic children (p=0.0001). This suggests 

cotinine exposure may be affected by asthma and GSTT1 or CYP2A6 polymorphisms.

183 Identification of Genomic Regions Linked to 

Epigallocatechin Gallate Induced Liver Toxicity Using the 

Diversity Outbred Stock.

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Epigallocatechin gallate (EGCG), an abundant polyphenol in green tea, has 

caused idiosyncratic liver toxicity when taken as an herbal supplement. The identi-

fication of genetic risk factors utilizing mouse population-based approaches, and 

validated in patient cohorts, could improve clinical management of EGCG-in-
duced liver injury. In order to map genomic loci related to EGCG induced hepato-

toxicity, we utilized the Diversity Outbred (DO) stock. DO mice are derived from 

eight inbred founder strains and have high genetic diversity, enabling high resolu-
tion mapping in this population. We hypothesized that Quantitative Trait Locus (QTL) 

mapping in DO mice exposed to EGCG would allow us to identify candidate 

genomic regions influencing the hepatotoxicity of EGCG. Male DO mice were 
treated once daily for 3 days, with EGCG (50 mg/kg i.p.) or vehicle. Twenty 

hour after the final dose, animals were sacrificed and serum and liver tissue 

were collected. Similar to humans, EGCG treatment in DO mice precipitates wide 

variation in hepatotoxic response. In treated animals, serum alanine aminotrans-

ferase (ALT) fold changes (terminal compared to pre-dose) ranged from 0.46-495.5 

(mean: 24.8 ± 65.4) and percent liver necrosis ranged from 0-86.8% (mean: 6.3 ± 14.1). 

QTL mapping in treated animals identified two suggestive loci— one on 

chromosome 12 and one on chromosome X. In a follow-up study, we will genotype 
suspected risk alleles in DNA collected by the Drug Induced Liver Injury Network 

(DILIN) from patients with suspected EGCG-induced liver toxicity. We have 

constructed the first application of the DO mice to the detection of xenobiotic 

risk alleles of toxicity responses. While further validation is needed, our data suggest 

that QTL mapping in DO mice may aid in identification of pharmacogenetic risk 

alleles for compounds causing liver injury.

184 Pharmacon-Genomics and -Genetics of 5-Fluouracil in 

Koreans.

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Genetic polymorphisms of several enzymes such as dehydroprimidine dehydroge-

nase (DHFR), thymidylate synthase (TS) have been emphasized for pharmacon-genomics and -genetics of 5-FU, which has been used for half century as a representative therapy for various cancers, however, shown individual variations in its various toxicities including life-threaten-
ing toxicity. Focusing on the three genes, we performed a pharmacon-genomic and 

-genetic study of 5-FU in a Korean population. Most of genotypes and gene ex-

pression were analyzed with 7500 Realtime PCR System (ABI). As results, we 

found genetic polymorphisms in DPDY-85, -1627, -1896 sites, 5’-ER, and 3’-

UTR at TS, and MTHFR-222, -227T, -237G sites among the Korean subjects (N=133; 

normal, N=109; head and neck patients, N=28). There was a significant association 

between 3’UTR genetic polymorphism at TS and its genotypic expression (p<0.05).

From 5-FU pharmacokinetic (PK) analyses, each genotype did not show any effect 

on PK parameters. However, the combination of genetic polymorphisms in 

MTHFR-222T/C, DPDY-1896T/C and 3’-UTR at TS showed significant differ-

ences in AUC of 5-FU and 5-FU/5-FUH2 ratios (p<0.01). Therefore, this study 

provides association between TS expression and its 3’-UTR polymorphism. 

Moreover, combination of the three genetic polymorphisms of the three genes can 

affect PK of 5-FU in Koreans.

185 PON1 Genotypes of Black Females from the Mississippi 

Delta Are Different from Those in the Rest of the State and 

Country.

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Paraoxonase (PON1) is named for its ability to hydrolyze paraoxon, the active 

metabolite of the insecticide parathion. It is implicated in cardiovascular and meta-

bolic diseases, such as Type 2 diabetes as well as tolerance of organophosphate in-

secticides. The human PON1 gene has several single nucleotide polymorphisms 

(SNPs). The SNP of an arginine (R) to glutamine (Q) substitution at codon 192 is 

associated with catalytic efficiency while the SNP of a methionine (M) to leucine 

(L) substitution at codon 55 is linked with serum levels. Since the Mississippi Delta 

has the highest rate of metabolic disease in the country, and a historically high 

level of pesticides, we compared the PON1 genotypes of black female Mississippi Delta 

clinic patients to those of black females from northeastern Mississippi and military 

populations. Genomic DNA was isolated from whole blood and the SNP codon 

areas were amplified by PCR. Since there is a native Alw I restriction site at codon 

192, the QQ, RR, and QR SNPs yield different digest patterns. A native NlaIII rest-

rection site at codon 55 generates different digest patterns for the LL, MM, and 

LM alleles. Genotypic frequencies were different for the Q192R SNP. QR was 

most frequent in the military (45%) and northeastern Mississippian (50%), but 

RR was most frequent in the Delta group (54%). Similarly, the most common 

combination of polymorphisms was QRLL in the military (24%) and northeastern 

Mississippi groups (36%), but in the Delta group it was RRLL (48%). Using 

Fishers exact test to analyze genotypic frequencies, the largest differences were be-

tween the Delta and military with Q192R P=0.0024 and L55M P=0.000176. When all 

groups were compared, Q192R ratios were significantly different at P=0.00007 

and L55M ratios at P= 0.00001. The significant differences seen in the Delta population may be associated with the region’s health dis-

parities.

186 In Vitro Toxicogenomic Screen Developed Using Genetically-

Diverse Mouse Inbred Cell Lines: Developing In Vitro 

Validations.

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Cell-based assays provide unprecedented means to globally and systematically 

screen for drugs and chemicals likely to display high inter subject toxicity variabil-

ity. Genetic determinants of toxicity identified in these screens can guide clinical 

trial design or identify susceptible subpopulations. Here, we developed an innova-

tive in vitro genetic screen using mouse embryonic fibroblast (MEFs) cells isolated 

from 32 inbred strains to screen them against 69 different drugs and chemicals.

Using high-content imaging, we measured multiplexed cell health parameters, in-

cluding cell loss, mitochondrial membrane potential, and cytochrome c release at 

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24 and 72 hours post treatment. We looked for genetic loci significantly linked to inter-strain cellular responses to treatment and found a 1.2 Mb locus on Chr X that was significantly linked to variable cytoxic responses to a known mitochondrial toxicant, rotenone (log2FDR > 4.0). Within this putative locus is cytochrome b-245, beta polypeptide (Cybb) gene, which encodes for a voltage-gated H+/K+ channel that mediates pH in the mitochondria. We conducted a series of experiments to examine the role of Cybb in mediating toxic responses to rotenone in vivo, given that mitochondrial dysfunction has been shown to underlie idiosyncratic adverse drug reactions. We found that strains belonging to different Cybb haplotypes scored differently in the treadmill exercise stress test for aerobic endurance after chronic treatment with rotenone. Our study demonstrates that cell-based genetic assays using MEFs are an effective tool for identifying genes underlying drug and chemical toxicity. Importantly, mouse strains that exhibit differential in vitro sensitivity, from a cell-based screen, also display a differential in vivo phenotype. This in vitro-to-in vivo validation is difficult, but a fundamental step toward recognition of cell-based toxicity screens.

(17%) than in controls (20%). The normal frequency of the GSTM1 negative genotypes was 52% in bladder cancer cases and in the controls as well and thus much lower, compared to a previous study performed from 1992-95 in the same area (70%). NAT2 genotypes were distributed equally among cases and controls (63% slow acetylators). Less GSTT1 negative genotypes were present in cases of bladder cancer compared to controls (OR 2.45, CI:1.65, 28.24), respectively. In conclusion, we replicated association between two closely linked CTNNA3 gene SNPs and DA in Caucasian workers. These findings suggest that genetically altered expression of CTNNA3 might influence cellular adhesion and the epithelial barrier function in the airways and play a role in the pathogenesis of DA.

A genome-wide association study conducted recently in Korean subjects identified three CTNNA3 (alpha-T catenin) single nucleotide polymorphisms (SNPs) (rs10762058, rs7088181, and rs4378283) associated with diisocyanate induced occupational asthma (DA). We conducted a candidate gene association study to replicate these findings in Canadian workers. Genotyping was performed on genomic DNA, using a 5’ nucleotide PCR assay. Genotyping of these SNPs was performed in 410 diisocyanate-exposed and predominantly Canadian workers including: 132 workers with DA confirmed by a specific inhalation challenge (DA+); 131 symptomatic workers in whom DA was excluded by a negative challenge (DA-); and 147 HDI-exposed asymptomatic workers (AWs). CTNNA3 rs7088181 and rs10762058 SNPs were significantly associated with DA+ when compared to AWs (p<0.05) but not in comparison to DA- workers. After adjusting for potentially confounding variables of age, smoking status and duration of exposure, minor allele homoygotes of rs7088181 and rs10762058 SNPs were at increased risk for DA compared with AWs. [OR= 9.05 (95% CI:1.60, 48.54) and OR = 6.82 (95% CI:1.65, 28.24), respectively]. In conclusion, we replicated association between two closely linked CTNNA3 gene SNPs and DA in Caucasian workers. These findings suggest that genetically altered expression of CTNNA3 might influence cellular adhesion and the epithelial barrier function in the airways and play a role in the pathogenesis of DA.

This work was supported in part by the NIEHS IAG (Y1-ES-0001) and NIOSH/CDC R01 OH 008795.
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Cytochrome P450 2S1 (CYP2S1) is one of the most recent additions to the P450 family of enzymes. Its expression is elevated in cancer and catalyzes the metabolic activation of the anticancer prodrug, AQ4N, under hypoxic conditions. Interestingly, our lab has also demonstrated that increased CYP2S1 expression may protect against AQ4 cytotoxicity under normoxic (21% O2) conditions. The main objective of this study is to determine whether individual variability in the CYP2S1 enzyme activity impacts the human lung cells to AQ4N and AQ4-mediated cytotoxicity. Five published CYP2S1 allelic variants have been published: CYP2S1*2 (R380C), CYP2S1*3 (P466L), CYP2S1*4 (S61N), CYP2S1*5 (L230R). According to the NCBI database, two additional non-synonymous variants have been identified in cancer (A205T and L189F) and L189F is restricted to African American populations. We generate each of the polymorphisms in a CYP2S1-Flag mammalian expression vector, using site directed mutagenesis. Thus far, two stable lines (S61N and L189F) in bronchial epithelial cells (BEAS-2B) and four stable lines (S61N, R166H, A205T, and P466L) alveolar carcinoma cell lines (A549) have been made. Examination of AQ4N and AQ4 cytotoxicity in BEAS-2B cell revealed that both mutants (S61N and L189F) exhibit significantly increased cytotoxicity in response to AQ4N compared with wild type CYP2S1-Flag and pDNA3.1 controls. Interestingly, the S61N polymorphism was significantly more sensitive to AQ4 than any of the other cell lines. This effect appears to be selective because cytotoxicity is not altered in response to a similar topoisomerase II inhibiting drug, etoposide. We are currently testing the effects of those polymorphisms on AQ4N and AQ44 levels, using HPLC. Research funded by NIH NIGMS Grant # R25GM061222.

Pancreatic cancer is the fourth leading cause of cancer deaths in the United States with a five year survival rate of less than 6%. Several environmental risk factors have been identified for pancreatic cancer, including dietary factors. In particular, high dietary intake of folate has been associated with a decreased incidence of pancreatic cancer, while low plasma folate levels are associated with an increased cancer risk. In addition, we found that SNPs in betaine hydroxymethyltransferase (BHMT; rs4027917) and folate levels and elevated homocysteine (Hcy) levels. In support of this, we found that cancer cases were more likely to express the allele associated with high BHMT activity, but not CMPA activity. Activity for either substrate was not found to be associated with CRP or TNF polymorphisms. BMI was found to significantly correlate with CMPA activity, but not CMPA. Activity for either substrate was not found to be associated with environmental exposure data suggested an association between pancreatic cancer and exposure to welding fumes. These findings suggest that increased BHMT activity may protect against pancreatic cancer risk.

Pancreas 1 (PON1) is an important HDL-associated endogenous antioxidant found to play a major role in susceptibility to health effects from pesticides and oxidant stress. A number of gene polymorphisms influence both protein concentration and substrate specificity, as do certain lifestyle factors. However, reports about the influence of PON1 genetics have varied widely in the literature. The goal of this study was to examine the influence of genes and health data in a cohort of monozigotic and dizygotic twins to better understand the effect of genetics on PON1 activity levels. DNA, serum, and standard blood donation information was obtained from 6 sets of dizygotic twins and 13 sets of monozygotic twins. DNA was genotyped for polymorphisms within PON1, PON2, C-reactive protein (CRP), and tumor necrosis factor (TNF). PON1 activity was determined using phenyl acetate (PA) and CMPA ([4-(Chloromethyl)phenyl acetate] as substrates, and genotyping was performed using the TaqMan Applied Biosystems 7900 HT System. Using a general linear model, PON1 Q192R, L55M and C-108T were significantly associated with both PA and CMPA activity, with CMPA activity showing a stronger association with genetic variants. PON2 S311C was significantly associated with PA activity, but not CMPA activity. Activity for either substrate was not found to be associated with CRP or TNF polymorphisms. BMI was found to significantly correlate with CMPA activity, but not CMPA. Activity for either substrate was not found to be associated with any environmental exposure data suggested an association between pancreatic cancer and exposure to welding fumes. These findings suggest that increased BHMT activity may protect against pancreatic cancer risk.

Environmental Health, Indiana University School of Public Health, Bloomington, IN.

Paraoxonase 1 (PON1) is an important HDL-associated endogenous antioxidant found to play a major role in susceptibility to health effects from pesticides and oxidant stress. A number of gene polymorphisms influence both protein concentration and substrate specificity, as do certain lifestyle factors. However, reports about the influence of PON1 genetics have varied widely in the literature. The goal of this study was to examine the influence of genes and health data in a cohort of monozigotic and dizygotic twins to better understand the effect of genetics on PON1 activity levels. DNA, serum, and standard blood donation information was obtained from 6 sets of dizygotic twins and 13 sets of monozygotic twins. DNA was genotyped for polymorphisms within PON1, PON2, C-reactive protein (CRP), and tumor necrosis factor (TNF). PON1 activity was determined using phenyl acetate (PA) and CMPA ([4-(Chloromethyl)phenyl acetate] as substrates, and genotyping was performed using the TaqMan Applied Biosystems 7900 HT System. Using a general linear model, PON1 Q192R, L55M and C-108T were significantly associated with both PA and CMPA activity, with CMPA activity showing a stronger association with genetic variants. PON2 S311C was significantly associated with PA activity, but not CMPA activity. Activity for either substrate was not found to be associated with CRP or TNF polymorphisms. BMI was found to significantly correlate with CMPA activity, but not CMPA. Activity for either substrate was not found to be associated with any environmental exposure data suggested an association between pancreatic cancer and exposure to welding fumes. These findings suggest that increased BHMT activity may protect against pancreatic cancer risk.

Pancreas 1 (PON1) is an important HDL-associated endogenous antioxidant found to play a major role in susceptibility to health effects from pesticides and oxidant stress. A number of gene polymorphisms influence both protein concentration and substrate specificity, as do certain lifestyle factors. However, reports about the influence of PON1 genetics have varied widely in the literature. The goal of this study was to examine the influence of genes and health data in a cohort of monozigotic and dizygotic twins to better understand the effect of genetics on PON1 activity levels. DNA, serum, and standard blood donation information was obtained from 6 sets of dizygotic twins and 13 sets of monozygotic twins. DNA was genotyped for polymorphisms within PON1, PON2, C-reactive protein (CRP), and tumor necrosis factor (TNF). PON1 activity was determined using phenyl acetate (PA) and CMPA ([4-(Chloromethyl)phenyl acetate] as substrates, and genotyping was performed using the TaqMan Applied Biosystems 7900 HT System. Using a general linear model, PON1 Q192R, L55M and C-108T were significantly associated with both PA and CMPA activity, with CMPA activity showing a stronger association with genetic variants. PON2 S311C was significantly associated with PA activity, but not CMPA activity. Activity for either substrate was not found to be associated with CRP or TNF polymorphisms. BMI was found to significantly correlate with CMPA activity, but not CMPA. Activity for either substrate was not found to be associated with any environmental exposure data suggested an association between pancreatic cancer and exposure to welding fumes. These findings suggest that increased BHMT activity may protect against pancreatic cancer risk.

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line and the effects of the haplotypes on MGMT transcription were determined using a direct competitive PCR assay. Compared with the most common (reference) haplotype, haplotypes 7 and 18 induced a significant 60% and 65% reduction in expression, respectively (P<0.001). However, haplotype 11 significantly increased expression by 70% (P<0.001). These data indicate that MGMT haplotypes regulate MGMT transcription and thus could play a major role in tumor response to TMZ treatment. Work is in progress to define the underlying mechanisms and to develop sensitive and specific markers that can distinguish those patients who would most be likely responsive to chemotheraphy from those who would not (supported by P30 ES006676; T32-07454; 1R03 NS065392-01 grants).

196 Associations between Genetic Polymorphisms of the Genes Mediating Inflammatory Response and Acute Pancreatitis Risk.

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Acute pancreatitis is a common inflammatory disease. The reported incidence is approximately 30 to 40 per 100,000 population per year and 25% will develop severe or life-threatening complications. Inflammation is typified by the activation of immunocytes such as monocytes and macrophages, and the secretion of inflammatory mediators such as nitric oxide, prostaglandin E2, and tumour necrosis factor-α (TNF-α). Nitric oxide is especially controlled by the inducible nitric oxide synthases (iNOS). Prostaglandin E2 is produced from arachidonic acid metabolites by the catalysis of cyclooxygenase-2 (COX-2). TNF-α, thought to be the first cytokine released, is a principal mediator of immune responses. Genetic factors may play important roles in susceptibility to pancreatic injury, as well as in the severity and evolution of the inflammatory process. The aim of our study was to determine if polymorphisms in iNOS, COX-2, TNF-α genes were associated with acute pancreatitis. For that, three iNOS (Ser608Leu, 1173C>T, 934G>C), seven COX-2 (rs5275, rs2206593, rs4684826, rs4684826, rs2066826, rs5277, rs2745577) and two TNF-α (308G/A, 238 G/A) variants were determined using polymerase chain reaction-restriction fragment length polymorphism analysis in patients with acute pancreatitis and healthy controls. Odds ratios (OR) and 95% confidence intervals (CI) were estimated. In conclusion; the association was seen with COX-2 rs5275 (P=0.03); specifically, patients carrying the TT genotype in comparison to patients carrying the CC genotype had a significantly lower risk of disease (OR=1.88; 95% CI:1.06-3.34). Both SNPs of TNF-α were not genetic risk factor for acute pancreatitis susceptibility. It was also found that iNOS Ser608Leu polymorphism was more frequent among cases with acute pancreatitis compared to controls (OR=2.88; 95%CI:1.49-5.57; P=0.002). We believe that the findings may be beneficial to the development of efficacious preventive strategies and therapies for inflammation-associated diseases.

197 In Vitro Toxicity of Antibacterial Silver Ions Released by Low Intensity Direct Electric Current (LIDC) Stimulation.

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Medical devices related surgical site infections and their treatment are a major cause of concern in the global healthcare system. The problem is compounded by the presence of antibiotic-resistant bacteria such as methicillin-resistant Staphylococcus aureus (MRSA) in healthcare environments, because infections caused by these bacteria are difficult to treat with conventional antibiotics. To prevent transmission of Staphylococcus aureus (MRSA) in healthcare environments, because infections caused by these bacteria are difficult to treat with conventional antibiotics. Two cytotoxicity tests commonly used in this evaluation are the MEM elution (ME) and Colony Formation (CF). The results from 43 samples used in medical devices were analyzed to evaluate the utility of both tests. Twenty-eight test samples also were evaluated for in vivo irritation, sensitization and systemic toxicity. The ME test was performed according to ISO 10993-3, section 8.2 including serial dilution of the extract, and CF test was performed according to ISO 10993-5. The in vivo tests were performed according to the appropriate ISO 10993 standard. The ME and CF test results from 36 test samples showed agreement (84% agreement). Seven test samples were non - toxic in ME but toxic in the CF test (16% disagreement). Toxicity was not observed in any of the in vivo tests on the 12 samples that were toxic in either the ME, CF, or both cytotoxicity tests. When compared to the results of 28 sets of in vivo tests, the ME test was 71 % in agreement and the CF test was 46% in agreement. Cytotoxicity was not observed in ME or CF tests for the 3 samples that failed for irritation. The results from this comparison demonstrate that in vitro cytotoxicity test is not predictive of in vivo irritation, sensitization or systemic toxicity; and the lack of cytotoxicity does not guarantee acceptable in vivo test results. The ME test with serial dilutions is more in agreement with results from in vivo tests, and when toxicity is observed provides comparable results to CF test.

198 Safety Assessment of Colorants Used in a Short Term Blood Contacting Medical Device—Challenges in Color Extraction Testing.


Use of colorants in medical devices continues to be scrutinized by the FDA. In response to the FDA questions on a recent PMA submission and to be compliant with ISO 10993 Biological Evaluation of Medical Devices - Part 1: Evaluation and Testing within a Risk Management Process, Boston Scientific completed extraction studies on colorants as part of the safety assessment. The challenges and successes in demonstrating the safety of the colorants used in a short-term blood-contacting catheter are presented here. A color elution study was conducted to demonstrate the potential bioavailability of four colorants. Devices were extracted at 37°C for 24 hours in aceton. These aggressive extraction conditions maximized the potential to extract colorant and were not intended to represent clinical use. Non-volatile residue (NVR) portions of the extracts were microwave digested and then analyzed using ICP-OES against control colorant samples of known concentration. The estimated maximum amount of extracted colorant ranged from <1.0 Q - 2.4 mg/de- vice. These values together with the NVR value (0.1 mg/device) from the USP Physicochemical test were used as ”worst case” estimates in the safety assessment. Variable process spike recovery results and device degradation during extraction contributed to uncertainty in the reported results. The safety assessment was conducted following ISO 10993-17: Establishment of Allowable Limits for Leachable Substances. The existing toxicological and biological safety data were included in the risk assessment. The threshold of toxicological concern (TTC) approach was applied for colorants with insufficient toxicological data. We concluded that the colorants used in this short-term catheter are eluted from the device at toxicologically insignificant amounts when extrapolated to the clinical exposure.

199 Comparison of Results from 2 In Vitro Cytotoxicity Tests Used in the Evaluation of Medical Devices.

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ISO 10993-1 includes an in vitro cytotoxicity test as part of a biological evaluation of medical devices. Two cytotoxicity tests commonly used in this evaluation are the MEM elution (ME) and Colony Formation (CF). The results from 43 samples used in medical devices were analyzed to evaluate the utility of both tests. Twenty-eight test samples also were evaluated for in vivo irritation, sensitization and systemic toxicity. The ME test was performed according to ISO 10993-3, section 8.2 including serial dilution of the extract, and CF test was performed according to ISO 10993-5. The in vivo tests were performed according to the appropriate ISO 10993 standard. The ME and CF test results from 36 test samples showed agreement (84% agreement). Seven test samples were non – toxic in ME but toxic in the CF test (16% disagreement). Toxicity was not observed in any of the in vivo tests on the 12 samples that were toxic in either the ME, CF, or both cytotoxicity tests. When compared to the results of 28 sets of in vivo tests, the ME test was 71 % in agreement and the CF test was 46% in agreement. Cytotoxicity was not observed in ME or CF tests for the 3 samples that failed for irritation. The results from this comparison demonstrate that in vitro cytotoxicity test is not predictive of in vivo irritation, sensitization or systemic toxicity; and the lack of cytotoxicity does not guarantee acceptable in vivo test results. The ME test with serial dilutions is more in agreement with results from in vivo tests, and when toxicity is observed provides comparable results to CF test.

200 Evaluation of Sample Preparation Methods in the ISO 10993-12 Standard: Implications for the Biocompatibility Assessment of Medical Devices.

R. P. Brown, H. Dinesdurage, J. Goode and M. Ghosh. US FDA, Silver Spring, MD.

The ISO 10993-12 standard outlines typical extraction conditions for the preparation of test samples for the biological evaluation of medical devices. The standard provides a list of temperature, time, and solvent conditions recommended for the preparation of extracts for toxicity testing, but no guidance is offered on which specific extraction conditions are optimal for various types of materials. The goal of this study is to identify extraction conditions that can be used to differentiate toxic from nontoxic polymeric materials in an indirect hemolysis assay without resulting...
in the degradation of the material. A ‘toxic’ material is defined in this study as one that produces a positive response in a modified MEM Elution cytotoxicity test. A wide range of polymeric materials (e.g., BUNA, nitrile, butyl, neoprene, latex, silicone rubbers; polyurethane, polyethylene, PVC) was extracted in a closed glass vial in phosphate buffered saline (PBS) using the default conditions outlined in the ISO 10993-12 standard (37°C x 24 hrs, 50°C x 72 hrs, 60°C x 72 hrs, 70°C x 24 hrs, 121°C x 1 hr). In addition, the polymers were extracted in 5% or 50% ethanol (EtOH) or acetone at 37°C for 24 hours. The 50% EtOH extracts were diluted to 5% with PBS for the hemolysis assay. The acetone extracts were evaporated under a nitrogen stream, then reconstituted with PBS. Our results show that rigorous extraction conditions (acetone, 50% EtOH) are necessary to correctly differentiate toxic from nontoxic materials. For example, latex and BUNA were positive in the cytotoxicity assay, positive when 50% EtOH extracts were used in the hemolysis assay, but negative when extracted in PBS using the standard extraction conditions in the ISO 10993-12 standard (e.g., 50°C x 72 hrs). Since the use of acetone as an extraction vehicle resulted in the degradation of some materials, such as PVC, the use of EtOH/PBS solvent mixture represents a promising approach for preparing samples for the biological evaluation of medical device materials.

**201 Can There Be a Universal Extraction Solvent for Medical Device Biocompatibility Testing? Comparison of Extraction Efficiencies among Five Solvents Used to Extract Polymeric Dental Devices.**

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The biocompatibility of medical devices is often evaluated using extracts prepared from the final product. The selection of the most appropriate extraction solvent(s) for producing chemical characterization and biocompatibility assessment remains a subject of active discussion within the medical device standards community. For this study, we examined extraction data obtained for nine experimental dental product prototypes extracted using five solvents of varying polarity to address both clinical use and exaggerated extraction scenarios. The extracted prototypes included four composite restoratives, a resin-modified glass ionomer, a polymeric polishing brush, a temporary cement, and a dental sealant. ISO 10993-12 compliant samples of each product were extracted in aqueous 5% ethanol solution, acetone, methanol, heptane, and 50-50 cyclohexane/isopropanol at 37°C with a target sample/solvent extraction ratio of 0.2 g/mL. Extraction time intervals ranged from 24 hours to 28 days. Gravimetric and HPLC analyses of the extracts show that in most cases use of methanol either resulted in the highest concentration of leachables or gave results similar to acetone. In the remaining cases, either acetone or water extracted the largest amount of residue. These results confirm the utility of methanol as an exaggerated solvent for many polymeric dental products, but also highlight the importance of proper solvent selection based on detailed knowledge of product chemistry.

**202 Determination of Total Leachable Bisphenol A from Polysulfone Membranes in Hemodialyzers and Hemoconcentrators.**

S. M. Cho, Y. Choi, H. Luu and J. Guo. US FDA, Silver Spring, MD.

Bisphenol A (BPA) is a high-production-volume chemical widely used in manufacture of polysulfone (PS), polycarbonate, epoxy resin, etc. Over the past ten years, BPA has been the subject of numerous risk assessment reviews and research worldwide because of its potential to produce adverse health effects through endocrine disruption. Although there is a significant body of literature focused on the adverse effects of BPA at low doses, there are discrepancies in the relevance and reliability of the published results. These make it difficult to properly evaluate the hazards of BPA. To reduce discrepancies and variation in research results, it is essential to establish reproducible/accurate analytical methods. In this study, we evaluated the BPA levels eluted from porous PS membranes used in hemodialyzers and hemoconcentrators using single and multiple consecutive extractions under clinically relevant extraction conditions. The levels of BPA release were determined using solid phase extraction (SPE) coupled with high performance liquid chromatography–mass spectrometry (HPLC-MS). We demonstrated that it was difficult to determine the total amount of BPA released from the PS membranes used in a single extraction method with finite solvent volume because of the chemical equilibrium between the extraction solution and the polymer phase. A general equation was derived to fit the BPA elution data and deepen our understanding on the equilibrium phenomenon during the extraction. The results revealed that repeated consecutive extractions of the PS membranes are needed to accurately determine the total leachable BPA in porous membranes.

**203 Bisphenol A Content in Polycarbonate from Medical, Automotive and Consumer Suppliers.**


Bisphenol A (BPA) is an organic compound used to make polycarbonate (PC) polymers and epoxy resins. The presence of BPA has the potential to produce human reproductive and developmental effects. Three sources of PCs were utilized in this study: from medical, automotive, and consumer suppliers (cups). Sterilized and unsterilized samples were extracted using ethanol (EtOH) or isopropanol (IPA) and incubated at 37 degrees C for 24 hours. Extracts were analyzed by high performance liquid chromatography (HPLC). In addition, exhaustive extraction by Soxhlet with IPA was performed on a medical grade PC. BPA was below the level of detection (0.5µg/g) in ETOH and IPA extracts with the exception of the cups. In extracts from cups, between 4 to 5.8µg BPA/g of test sample was detected, however BPA was not detected in the non-sterilized sample extracted with ETOH (limit of detection was 0.5µg/g). Assuming a 10 cup and the worst-case of 5.8µg BPA/g, an adult male (70kg) would be exposed to 0.82µg BPA/kg/day. Using exhaustive Soxhlet extraction with IPA, BPA was below the limit of detection (0.5µg/g) for automotive PC. Assuming a 10 g sample and the worst-case of 0.5µg/g, an adult male would be exposed to 0.071µg/kg/day. The average BPA amount found in medical grade PC by exhaustive Soxhlet extraction with IPA was 0.25µg/g. For a 10g medical device, the calculated exposure for an adult male is 0.036µg/kg/day. The US FDA acceptable Daily Intake (ADI) and/or EU Tolerable Daily Intake (TDI) for BPA is 50µg/kg/day. The average adult male exposures to BPA from consumer, automotive, and medical grade PC are 58, 704, and 1300 times less than US FDA ADI or EU TDI. The extractable BPA from automotive and medical grade PC was significantly less than that observed from the consumer PC.

PS

**204 Local Effects of Microelectrode Implantation in Rabbit Muscles.**

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The purpose of this study was to evaluate the biocompatibility of various polymer-based microelectrodes (PBMs) after implantation in rabbit muscle tissues following a standardized method. Three types of PBMs were examined: silicone-based platinum, polimide-based gold, and liquid crystal polymer-based gold microelectrodes. All experimental procedures followed the International Organization for Standardization (ISO) 10993-6:2007(E). Six female rabbits were used for this study. The PBMs were implanted into the left paravertebral muscle of the dorsal region of the rabbits for 12 weeks, each type being implanted into two rabbits. Control article (high density polyethylene, HDPE) was implanted in the equivalent site on the right side of each rabbit. No changes in the clinical signs, mortality, body weight, and gross findings related to the PBMs were noted. The results of histopathological evaluation suggest that the PBMs did not induce any cellular changes. Thus it could be concluded that the three types of PBMs are all non-toxic, non-irritating, and biocompatible.

**205 Subchronic Systemic Toxicity of Subcutaneous Implantation of Microelectrodes in Rats.**

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The purpose of this study was to evaluate the biocompatibility of various polymer-based microelectrodes (PBMs) through the subchronic systemic toxicity of subcutaneous microelectrode implantation in rats following a standardized method. Three types of PBMs were examined: silicone-based platinum, polimide-based gold, and liquid crystal polymer-based gold microelectrodes. All experimental procedures followed the International Organization for Standardization (ISO) 10993-6:2007(E) and ISO 10993-11:2006(E). Ten female rats were used for four each groups. Control article (high density polyethylene, HDPE) and three types of PBMs were implanted subcutaneously in the same site in each group and were left in place for 13-weeks. No effects related to the PBMs were observed in any tested criteria, included mortality, clinical signs, body weight, food and water consumption, hematology and serum biochemistry parameters, urinalysis and ophthalmoscopy.
organ weight, gross findings, or histopathological findings. These results suggest that no subchronic systemic toxicity is induced by subcutaneous implantation of these three types of PBMs under the conditions used in this study.

206 Comprehensive Health-Based Risk Assessment of Material from an Ingestible Medical Device.
M. A. Nascarella1, G. M. Savage2, G. Moon2 and B. D. Beck1
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We present an analysis of the potential toxicity of an ingestible medical device. The primary toxicological concern is an 8 μm layer of copper (Cu), with an area of approximately 1.0 mm², that is a component of the device's battery. The Cu content is approximately 20-33.1 μg/device. We calculated the potential toxicological risks of Cu leached from the device, assuming a maximum use of 30 devices/event. Depending upon fluid in the stomach, an individual could have a stomach dose of approximately 219 μg Cu, resulting in a stomach concentration ranging from 0.25-2.73 μg Cu/mL. These concentrations may be compared to a threshold Cu concentration for gastrointestinal (GI) toxicity of 1.4 μg/mL. In the most plausible scenario, the predicted concentration of Cu in the stomach is below the concentration associated with GI symptoms in humans, consisting of mild, reversible effects and no associated systemic toxicity. The higher potential stomach concentrations somehow exceed the threshold concentration, but potential GI symptoms could be mitigated by ingesting the devices with food. We also estimated the potential total intake of Cu from all sources, including the device. Ingestion of the device, combined with the ingestion of median levels of Cu in food, water, and multivitamins, is estimated to be well below the 10 mg/day IOM determined safe daily intake for the general population. We also evaluated the cytotoxicity of the extractable material from this device, based on ISO-compliant tests of device extractions using simulated gastric fluid. Using open-source software, we calculated the number of devices that would be associated with a cytotoxic effect according to ISO standards. This analysis indicated that plausible use of the device would not lead to cytotoxic effects. Overall, we conclude that ingestion of the medical device under plausible use conditions is unlikely to present a toxicological concern for Cu.

207 Comparative Pulmonary Response to Aerosolized Humidifier Disinfectants by Intratracheal Instillation and Inhalation Exposure.
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1Korea Institute of Toxicology, Daejeon, Republic of Korea; 2Jeonbuk Department of Non-Human Primate, Korea Institute of Toxicology, Jeongeup, Republic of Korea.

Sponsor: S. Park

Mice intratracheal instillation (IT) and rat inhalation exposure (IE) were conducted to identify the toxicity of 3 representative humidifier disinfectants (products A, B, and C) containing polyhexamethylene guanidine (PHMG), 5-chloro-2-methylpyridine (5-CPM), and 5-chloro-2-methylpyridine-2-ethyl(2-ethyl)-2-chloro-5-bromo(thiol)-2H-oxazoline (HCBO). In mice administered by multiple 7-9 IT at 0.05 ml for 2 wk, severe necrotic obliterative bronchiolitis (OB) was found in product A and C. But no adverse treatment effect was observed in product B. In rat inhalation study (IE), necrotizing inflammation was observed in nasal and larynx, trachea, and bronchi of rat with product A at 0.4 mg/ml and product C at 1.75 mg/ml. However, necrotizing inflammatory lesions in the upper airways were not present in the IT due to direct administration of test substance to the lung via trachea. Granulomatous OB, bronchitis, collagenized fibrosis, alveolar bronchiolization, and extensive squamous metaplasia were observed in product A at 10 wk IH, and product C at 7 wk IH. No treatment-related adverse effects were observed in 13 wk IH with product B at 1.80 mg/ml. Lung lesions induced by IT and IH with product A and C were comparable and no treatment-related lesions were present with product B in both IT and IH exposure. It was difficult to evaluate dose-related toxicity by IT dosing. However, IT with low dose was a useful methodology to screen and identify toxicity of test substances in this study.

208 Effect of Fuel Composition on Chemistry and Pulmonary Toxicity in Mice Exposed to Biomass Pyrolysis Vapor.
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1NIEERL, US EPA, Durham, NC; 2NRML, US EPA, Durham, NC; 3CEMALB, University of North Carolina at Chapel Hill, Chapel Hill, NC.

Biomass pyrolysis is a method to form oil through the thermal degradation of organic material in the absence of oxygen. The process results in a mix of gaseous oxides, organic vapors and particulate matter that could pose an inhalation hazard. In this study, we characterized aerosol emissions from three different fuels (corn, pine, maple) at different conditions, compared the resultant system toxicity endpoints in CD-1 mice after a 4h inhalation exposure. Particle number counts/cc were 37000, 48000 and 34000 for corn cob, pine and maple respectively, with median count diameters of 221, 254 and 225 nm. Particle mass was 409, 689 and 362 μg/m3 for the three fuels, and CO measurements were 37, 39 and 45 ppm respectively. Particle size analysis by dynamic light scattering analysis of the three fuels showed acrolein being 1.3, 1.7 and 2.3 μm, with lower levels of propylene, acetone and vinyl acetate. Pulmonary responses were assessed in a plethysmograph immediately before and after exposure, as well as 4 and 24h post-exposure, when mice were euthanized. IL-6 and IL-10 levels were measured immediately following exposure, and this effect persisted at the 4h time point for the corn cob and maple atmospheres before returning to control levels at 24h. Total protein was increased in the BALF of animals exposed to corn cob (4h and 24h post), pine (24h post), and maple (4h post), with corn having the highest effect. No other endpoints were affected except BALF LDH for pine, and increased hematocrits for corn cob at 24h. Because these atmospheres were considered to have high irritant characteristics, nasal lavage was also performed and although some increases in inflammatory cells were seen for the corn cob and maple atmospheres, the effects were variable. We conclude that corn cob and maple pyrolysis products seemed to have a more potent effect on pulmonary function and toxicity parameters than pine emissions. (This abstract does not reflect EPA policy).

209 Gene Expression in Bronchiolitis Obliterans-Like Lesions in Rats Exposed to 2, 3-Pentanediol.
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Obliterative bronchiolitis (OB) is an irreversible lung disease characterized by progressive fibrosis in the small airways with eventual obliteration of the airway lumens. OB is most commonly associated with lung transplant rejection; however, OB has also been diagnosed in workers exposed to artificial butter flavoring (ABF) vapors. Research has been limited by the lack of an adequate animal model of OB, and as a result the mechanism is unclear and there are no effective treatments for this condition. A rat model of chemical-induced OB using the ABF component, 2,3-pentanediol (PD), was found to cause airway lesions histopathologically similar to OB lesions in humans. We used this model to evaluate changes in gene expression in the distal bronchi of rats with OB. Male Wistar Han rats were exposed to 200 ppm PD or air (controls) 6hr/d, 5d/wk for 2-wks. Distal bronchial tissues were laser microdissected from serial sections of frozen lung. In exposed lungs, both fibrotic and nonfibrotic airways were collected. Following RNA extraction and microarray analysis, differential gene expression was evaluated. In exposed nonfibrotic bronchi, 1548 genes were significantly altered relative to air-exposed controls with notable downregulation of many inflammatory cytokines and chemokines. In contrast, PD-exposed bronchi showed significantly altered with a majority of genes being upregulated in affected pathways. TGF-beta2 and downstream genes implicated in fibrosis were significantly upregulated in fibrotic lesions. Genes for collagens and extracellular matrix proteins were highly upregulated. In addition, expression of genes for peptides and for peptide inhibitors were significantly altered suggesting tissue remodeling that may contribute to fibrosis. These data will be used to gain a better understanding of the molecular mechanisms of OB and to identify potential therapeutic targets.

210 Polyhexamethylene guanidine Phosphate Induces Severe Lung Inflammation, Fibrosis, and Thymic Atrophy.
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Polyhexamethylene guanidine phosphate (PHMG-ph) has been widely used as a disinfectant due to its strong bactericidal activity. But the Korea Centers for Disease Control and Prevention (KCDC) and Ministry of Health and Welfare reported that humidifier disinfectants might cause of the unknown pulmonary disease in 2011. The purpose of this study was to assess the potential adverse effect of PHMG-ph, a ingredient of humidifier disinfectant, exposed to lung directly. 0.0125%, 0.0375%, and 0.0625% PHMG-ph was instilled intratracheally into CD-1 mice. Seven and fourteen days after instillation, lungs were collected and proinflammatory cytokines, chemokines and fibrotic markers were measured from lung lysates. We also performed flow cytometry to evaluate the cell distribution of thymus and RT-PCR to measure the mRNA expression associated with T cell development.
As a result, single exposure of 0.0125%, 0.0375%, and 0.0625% PHMG-Ph induced inflammatory response with increased neutrophils, macrophages, and immune cell infiltration to the lungs, and interestingly, this inflammation did not resolve till the end of the experiments (14 days after instillation). The histopathology showed the both inflammation and pulmonary fibrosis exacerbated at day 14 after exposure in dose-dependent manner. Also PHMG-Ph decreased the total cell number and the CD4+/CD8+ cell proportion in thymus and induced severe medulla reduction based on histopathology data. These observations demonstrated that PHMG-Ph exposed to lung lead to pulmonary inflammation and fibrosis as well as thymic atrophy.

211 Inhalation of a Spot Welding Aerosol Using an Adhesive Increased Airway Resistance but not Lung Inflammation.


Spot welding (SW) is used in the automotive and aircraft industries where high speed repetitive welding is needed and relatively thin metal sections are welded. Epoxy adhesives are applied as sealers to the seams of the metals that are joined. SW produces complex aerosols composed of both metal and volatile compounds which may cause bronchitis and asthma in workers. The goal was to assess the effect of SW fumes on lung function and toxicity. Male Sprague-Dawley rats were exposed by inhalation to 20 mg/m³ of SW aerosol in the presence of an adhesive for 4 hr/s x 8 d. Controls were exposed to air. Size distribution of the aerosol as determined by a MOUDI particle impacter was tri-modal with a MMAD of 1.6 μm in the large-size mode, 0.30 μm in the small-size mode, and 0.01-0.05 μm in the ultralfine mode. Two distinct particle morphologies were observed: a brownish metal particle that predominated in the small-size fraction and a black, glue-like particle that was in the large-size fraction. The metal fraction was found to be >90% Fe. Significant amounts of volatiles (e.g., benzene, toluene, others) were present, likely produced from the vaporization of the adhesive. At different times after exposure, bronchoalveolar lavage (BAL) was performed to assess lung toxicity. Lung resistance (Rl) was evaluated in a separate set of animals before and after challenge with inhaled methacholine (MCh). Immediately after exposure, baseline Rl was significantly elevated in the group exposed to the SW fumes. Basal Rl returned to control levels by 1 d after exposure. Reactivity to MCh was not affected at any time point after fume exposure. No significant increase in lung inflammation (neutrophil influx) or injury (cytotoxicity and lung epithelial permeability) was observed in BAL fluid at 1 and 5 d after exposure to SW fumes. Acute inhalation of SW fumes at occupationally-relevant concentrations may act as an irritant as evidenced by the increased Rl, but had little effect on toxicity.

212 Cardiopulmonary Health Effects of Traffic-Related Air Pollutants in a Healthy Population.

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There is emerging evidence that inhaling certain components of ambient particulate matter, specifically traffic pollutants, is associated with adverse health effects. We hypothesized that exposure to air pollution components of diesel exhaust-rich traffic, compared to cars-only traffic, produces greater adverse cardiopulmonary effects. In this case-crossover study, 23 participants were recruited to measure pulmonary function, exhaled NO, blood cytokines, heart rate variability, and blood pressure prior to, immediately after, and 24 hours after intermitting walking along 3 diverse roadways. Exposures lasted for 1.5 hours between June and September in 2011 and 2012, and personal exposures to pollutants were collected. The 3 locations differed by traffic type: the George Washington Bridge (GWB) carries truck and car traffic, the Garden State Parkway (GSP) carries only car traffic, and Sterling Forest, NY (SF) acted as a control location. Levels of PM2.5, PM10, black carbon, elemental carbon, and organic carbon were found to be highest at GWB and lowest at SF for all pollutants measured. The traffic count was similar between GSP and GWB. Using a repeated measures 2-way ANOVA, p-values were generated for time, location, and interactions between time and location. Cytokines was a significant factor for FVC (p = 0.04) and FEV1 (p = 0.05); a significant interaction term for pulse pressure was also observed (p < 0.01). Upon further analysis, systolic and pulse pressures varied significantly amongst locations when comparing the baseline and 24 hr-post-measurement, while IL-1b varied amongst locations between the baseline and immediately after exposure. A trend of increasing sNO at the GWB was seen immediately after exposure, but did not reach significance (p = 0.06). These results suggest that acute effects of traffic-related pollution are observed in a small, healthy population; these effects differed by traffic type.

213 Ventricular Transcriptional Data Provide Mechanistic Insights into Diesel Exhaust-Induced Attenuation of Cardiac Contractile Response and Blood Pressure.


Human exposure to diesel exhaust (DE) has been associated with cardiovascular impairments however the mechanisms and the role of hypertension are not well understood. We have shown that DE reduces blood pressure (BP) and cardiac contractility in healthy normotensive Wistar Kyoto (WKY) rats. We hypothesized that DE would induce differential myocardial gene expression changes that modulate contractility in WKY and spontaneously hypertensive (SH) rats, and that lowering BP in WKY and SH with hydralazine (HYD) would increase this effect of DE. Male WKY and SH rats were treated with HYD (150 mg/L) in drinking water for 10 days prior to exposure and until necropsy. All rats were exposed to clean air or freshly-generated whole DE (1500 μg/m3), 5-hrs/day for 2 days. Systolic BP was monitored using the tail-cuff method on days -10, 0, and 2. Left ventricular genome-wide expression was analyzed using Illumina RatRef-12 BeadChips. As expected, WKY and SH rat's ventricular gene expression patterns differed markedly. Surprisingly, DE exposure caused differential expression of 256 genes in WKY but not in SH rats. In WKY rats, the effect of HYD on expression patterns were nearly identical to changes induced by DE (same genes with same directional change); while HYD was without effect on expression changes in SH rats despite lowering BP. Genes inhibited by DE or HYD in WKY were related to decreasing BP and muscle contraction as well as calcium homeostasis and apoptosis. In conclusion, acute DE exposure caused gene expression changes only in normotensive WKY rats; these changes mimicked those induced by HYD and are associated with decreased cardiac contractility and BP in healthy rats. (Abstract does not reflect USEPA policy).

214 Comparative Cardiopulmonary Toxicity of Soy Biofuel and Diesel Exhausts in Healthy and Hypertensive Rats.

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Increased use of renewable energy sources raise concern about health effects of emissions from such sources. We conducted a comprehensive analysis of relative cardiopulmonary health effects of exhausts from 1) 100% soy biofuel (B100), 2) 20% soy biofuel + 80% low sulfur petroleum diesel (B20), and 3) 100% petroleum diesel (B0) in rats. Normotensive Wistar Kyoto and spontaneously hypertensive rats were exposed to these 3 exhausts at 0, 50, 150 and 500 μg/m3, 4 h/day for either 2d or 4 wk (5 d/wk) to mimic near environmental concentrations. Additionally, WKY rats were exposed for 1d and responses were analyzed 0 h, 1d or 4d later for time course analysis. Hematological parameters, in vitro platelet aggregation, bronchoalveolar lavage fluid (BALF) markers of pulmonary injury and inflammation, ex-vivo aortic ring constriction, heart and aorta mRNA markers of atherogenesis, and serum biomarkers of acute cardiac injury as well as cytokines were analyzed. The presence of macrophagedalons in the lung alveoli was clearly evident with all 3 exhaust exposures. Overall, exposure to all 3 exhausts produced only modest effects in most endpoints analyzed in both rat strains. B20 γ-glutamy transferase (GGT) activity was the most consistent marker shown to be increased in both strains with all 3 fuels (B0-B100-B20) without increases in B20 neoptilol. Small inconsistent changes in aorta mRNA markers of inflammation, vasoconstriction and thrombosis, and those of serum biomarkers need to be interpreted cautiously. Our comparative evaluations show modest cardiovascular and pulmonary effects at low concentrations of all exhausts. Additionally, our study highlights the value of BALF levels of GGT activity as the most sensitive biomarker in low level inhalation studies. (This abstract does not represent USEPA policy).
Acute and Delayed Effects of Intermittent Ozone on Cardiovascular and Thermoregulatory Responses of Young and Aged Rats.

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Ozone (O3) is associated with cardiovascular and respiratory diseases. The aged population is considered to be more sensitive to air pollutants but relatively few studies have demonstrated increased susceptibility in animal models of aging. To study the acute and delayed physiological responses to O3, core temperature (Tc) and heart rate (HR) monitored by telemetry in adult (12 m) and senescent (24 m) Brown Norway rats exposed to intermittent O3 (1.0 ppm, 6 hr/day for 2 consecutive d for 12 wk). Tc and HR dropped precipitously in both age groups during the 1st bout of O3 exposure; Tc decreased from -38 to -35 °C while HR decreased from -360 to -175 b/min. These acute responses were attenuated during the 2nd day of O3. As O3 exposure progressed, the acute Tc and HR responses abated but the aged animals were consistently less affected than the young adults throughout the 12 wk exposure period. During 5 d of recovery in home cages, both young and senescent rats displayed a fever-like -0.5 °C elevation in daytime Tc. HR was also elevated in the young adults during recovery. The rise in Tc persisted for 2-3 d after O3. As O3 exposure progressed, the acute Tc and HR responses abated but the aged animals were consistently less affected than the young adults throughout the 12 wk exposure period. During 5 d of recovery in home cages, both young and senescent rats displayed a fever-like -0.5 °C elevation in daytime Tc. HR was also elevated in the young adults during recovery. The rise in Tc persisted for 2-3 d after O3. As O3 exposure progressed, the acute Tc and HR responses abated but the aged animals were consistently less affected than the young adults throughout the 12 wk exposure period. During 5 d of recovery in home cages, both young and senescent rats displayed a fever-like -0.5 °C elevation in daytime Tc. HR was also elevated in the young adults during recovery. The rise in Tc persisted for 2-3 d after O3. As O3 exposure progressed, the acute Tc and HR responses abated but the aged animals were consistently less affected than the young adults throughout the 12 wk exposure period.

Rationale: Human controlled exposure studies have generally focused on subjects exposed to ozone (O3) while exercising. We exposed resting subjects to labeled O3 (nvHC) was generated under a modified Canadian Intense Regimen (CIR) and its biological activities were compared to those of Reference (3R4F) cigarette, using nose-only inhalation studies. In a 5-wk inhalation study, female SD rats were exposed to MS of either cigarette at 600 or 1000 μg wet total particulate matter (WTPM)/L for 1 hr, 2 times/day, 7 days/week for 5 weeks. Pulmonary inflammation was significantly weaker in nvHC groups compared to 3R4F groups, based on the neutrophil counts and deviation enzyme levels in bronchoalveolar lavage fluid (BALF). After a 4-week recovery, BALF parameters of nvHC groups were similar to the air-exposed Sham group, while those of 3R4F groups remained elevated. In a 13-wk inhalation study, male and female SD rats were exposed to MS from each cigarette at 200, 600, or 1000 WTPM μg/L for 1 hr/day, 7 days/week for 13 weeks. Histopathological changes in the respiratory tract were significantly lower in incidence/severity for nvHC groups, especially in respiratory epithelial hyperplasia and accumulation of pigmented macrophages in alveoli. After a 13-wk recovery, the lesions were completely or partially regressed, except for accumulation of pigmented macrophages in alveoli, in both nvHC and 3R4F groups. In conclusion, nvHC demonstrated clearly and significantly lower biological activities compared to 3R4F, based on the BALF parameters and histopathology.

Dose and Effect of Inhaled Ozone in Resting versus Exercising Human Subjects: Comparison with Resting Rats.


Methods: We measured O3 dose as the concentration of 16O in cells and extracellular material of nasal, bronchial and bronchoalveolar lavage fluid (BALF) immediately post exposure and related these measurements to O3 effects on inflammation, epithelial permeability and phagocytosis in the same fluids and to breathing parameters measured during the 18O3 exposure. A parallel study of resting subjects examined cytokine changes during and immediately following a 2 hr exposure to 0.18, 0.25, 0.3 and 0.4 ppm O3.

Results: Subjects exposed while resting had 18O concentrations in BALF and nasal lavage that were proportional to the amount of air breathed during exposure. Significant small changes were observed in BALF total cells and neutrophils and in BALF cell phagocytosis following resting O3, however, most indicators of O3 effects that were observable in exercising subjects (including increased BALF super-natant protein, lactate dehydrogenase, interleukin-6 and low molecular weight antioxidants) were not observed in resting subjects. The 18O incorporation into BALF of resting humans was similar to that of similarly exposed resting F344 rats. FEV1 changes in resting human subjects showed a much attenuated response compared to exercising subjects.

Conclusions: Quantitative measures of alveolar O3 dose and toxicity that were observed previously in exercising subjects were greatly reduced or non-observable in O3 exposed resting subjects. Resting rats and resting humans have similar alveolar O3 dose. Disclaimer: This abstract does not represent EPA policy.

Biological Responses in Rats Exposed to Mainstream Smoke from a Heated Cigarette Compared to a Conventional Reference Cigarette.

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The heated cigarette (HC) generates mainstream smoke (MS) primarily by vaporizing the components of the tobacco rod using a carbon heat source at the cigarette tip. Consequently, MS from HC contains markedly less chemical constituents compared to conventional (combusted) cigarettes. In this study, MS from a non-ventilated HC (nvHC) was generated under a modified Canadian Intense Regimen (CIR) and its biological activities were compared to those of Reference (3R4F) cigarette, using nose-only inhalation studies. In a 5-wk inhalation study, female SD rats were exposed to MS of either cigarette at 600 or 1000 μg wet total particulate matter (WTPM)/L for 1 hr, 2 times/day, 7 days/week for 5 weeks. Pulmonary inflammation was significantly weaker in nvHC groups compared to 3R4F groups, based on the neutrophil counts and deviation enzyme levels in bronchoalveolar lavage fluid (BALF). After a 4-week recovery, BALF parameters of nvHC groups were similar to the air-exposed Sham group, while those of 3R4F groups remained elevated. In a 13-wk inhalation study, male and female SD rats were exposed to MS from each cigarette at 200, 600, or 1000 WTPM μg/L for 1 hr/day, 7 days/week for 13 weeks. Histopathological changes in the respiratory tract were significantly lower in incidence/severity for nvHC groups, especially in respiratory epithelial hyperplasia and accumulation of pigmented macrophages in alveoli. After a 13-wk recovery, the lesions were completely or partially regressed, except for accumulation of pigmented macrophages in alveoli, in both nvHC and 3R4F groups. In conclusion, nvHC demonstrated clearly and significantly lower biological activities compared to 3R4F, based on the BALF parameters and histopathology.

A Cross-Regulatory T Cell Response in Pulmonary Hypertension.

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Exposure to urban air pollution (fuel emissions, particulate matter) has been associated with the exacerbation of autoimmune diseases. Our studies are focused on the mechanism of immune response induced pulmonary hypertension. We have shown that co-exposure of mice to inhaled antigen and urban particulate matter (PM) exacerbates pulmonary arterial remodeling and induces pulmonary hypertension. The current studies were performed with neutralizing anti-cytokine antibodies to identify the critical mediators for pulmonary hypertension and the interactions in the mediator network.

Sensitized mice were intranasally challenged with either antigen (Ovalbumin) combined with urban PM2.5 (collected in New York City), or given saline. Groups of mice were injected with neutralizing anti-interleukin (IL)-13, or anti-IL-17A/F antibodies alone or in combination, or control antibody. Right ventricular systolic pressures and immune response markers in the lungs were measured. Intranasal challenge with antigen and urban PM significantly increased right ventricular systolic pressures. Only combined, but not single, injections with IL-13- and IL-17A/F-blockers significantly reduced this outcome. Surprisingly, injections with single neutralizing antibodies not only significantly reduced the inflammatory markers known to be regulated by IL-13 or IL-17A/F, but also revealed cross-inhibition of these markers. For example, the increased expression of the antigen presentation molecule, major histocompatibility class II (MHCII), by airway dendritic cells was inhibited by the IL-13 blocker given alone or in combination, while the IL-17A/F blocker led to an increase in MHCII expression. Conversely, infiltration of the airways with neutrophils was inhibited by the administration of the IL-17A/F blocker given alone or in combination, while injections with the IL-13 blocker increased neutrophil infiltration. In conclusion, exposure to antigen and urban PM induced pulmonary hypertension by elaborating a mixed immune response that has at least two, cross-regulatory arms that are controlled by IL-13 and IL-17A/F, respectively.

Capsule-Based Aerosol Generator (CBAG)—Validation in a Rat Model of LPS-Induced Nonallergic Pulmonary Inflammation.


Intratracheal (IT) insufflation is the principal method of delivery of inhaled drug substances to conscious non-clinical species in early drug development; however, this achieves particulate deposition dissimilar to conscious inhaled delivery and can produce artefactual toxicological and pharmacological results. The CBAG was developed(1) as an alternative to IT insufflation whilst providing representative inhalation exposure by demonstrating the effectiveness of the CBAG in the rat model of LPS-induced non-allergic airway inflammation. Rats were exposed to 0.01, 0.1 or 1.0 mg/kg of inhaled fluticasone propionate (FP) over a 20-minute period using nominally 1 mg filled hydroxypropyl methyl cellulose size 2 capsules at blend strengths of 1, 10 and 100% w/w of FP respectively. Two concurrent control groups were exposed to lactose only using the same regime. Twenty minutes after the end of the inhalation exposure the animals were challenged with either aerosolised LPS (0.1 mg/mL) for the FP groups and one control group or 0.9% w/v saline (second control group) for 30 minutes. Rats were euthanized 4hrs following the challenge
and a bronchoalveolar lavage (BAL) investigation performed. A BAL total and differential cell count was used to evaluate the efficacy of PI. Delivered doses of 0.0103, 0.117 and 0.863 mg/kg were achieved, which were within 14% of target. This resulted in a dose dependent inhibition of BAL neutrophils of 40%, 79% and 98% respectively compared with the lactose/LPS control group. In conclusion, the results give confidence that the CBAG is a viable alternative to IT methodology for protecting lung health.


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### 220 Distinct Inflammatory Macrophage Subpopulations and Myeloid-Derived Suppressor Cells Accumulate in the Lung and Spleen following Exposure of Mice to Inhaled Ozone.

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Ozone is an ubiquitous urban air pollutant known to damage the lung. Activated macrophages (MP) and inflammatory mediators they have been implicated in ozone toxicity. However, the phenotype and origin of these cells have not been established. In these studies, techniques in flow cytometry were used to assess macrophage subpopulations in the lung, spleen and bone marrow following ozone inhalation. Exposure of C57BL/6 male mice to ozone (0.8 ppm, 3 h) resulted in increased bronchoalveolar lavage (BAL) protein levels after 24-72 h, indicative of alveolar epithelial injury. This was correlated with a rapid and persistent increase in the percentage of CD11b+F4/80+ inflammatory macrophages in BAL. An increase in F4/80 negative CD11b+Ly6C+Ly6G+ myeloid-derived suppressor cells (MDSCs) was also observed in BAL, a response most prominent 24 h post ozone exposure. Conversely, F4/80 positive CD11b+Ly6C+Ly6G+ MDSCs decreased in BAL after ozone exposure. We also found that ozone exposure resulted in a persistent decrease in CD11b+F4/80+ inflammatory macrophages, and a transient increase in CD11b+F4/80+Ly6C+Ly6G+ MDSCs in the spleen. In contrast, there were no changes in bone marrow cell subpopulations after ozone inhalation. Taken together, these results suggest that the spleen is a source of inflammatory MP in the lung following ozone exposure; moreover, subpopulations of MDSCs originating in the lung and the spleen may contribute to early inflammatory responses in the lung and to processes of injury and repair. Supported by NIH grants GM034310, ES004738, CA132624, AR055073, ES007148, ES005022.

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### 221 Compare In Vitro Endothelial Cell Release of Endothelium Derived Vasodilators in Response to Diesel, Biodiesel Blend and Biodiesel Neat Combustion Extracted.

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Diesel exhaust exposure in controlled human chamber studies found exposure induced inhibition of vasodilation. Particles emitted in exhaust can translocate into the vascular system however when particles are dissolved in solvents of various polarity the insoluble fraction separates from the soluble fraction. We collected the soluble fraction of combusted particles dissolved in DMSO for evaluation of the extract to interfere with the release of endothelium derived vasodilators. Endothelium dependent vasodilation is dependent on 6-keto PGE1 alpha (6keto) a vaso-active metabolite of arachidonic acid. We have investigated the effects of diesel, biodiesel blend and biodiesel neat for a change in 6keto release. QPCR data from extract exposure indicates there is no increase in markers of inflammation. However there is a measurable increase in heme oxygenase-1 (HO-1) gene expression. HUVEC and HCAEC exposed to extract for 6, 8 and 24h indicate no statistically significant change in 6keto release. QPCR data from extract exposure indicates there is no increase in markers of inflammation. However there is a measurable increase in heme oxygenase-1 (HO-1) gene expression. HUVEC and HCAEC exposed to 8h extract exposure to B100 at 100μg/mL have over two fold increase in HO-1. The B100 particle composition analysis indicates high levels of Zn and Fe compared to biodiesel blend and diesel. In our work we address a possible mechanism for attenuation of vaso-active arachidonic acid metabolites in endothelial cells exposed to diesel, biodiesel blend and biodiesel neat particle extracts. [This is an abstract of a proposed presentation and may not necessarily reflect official US EPA Policy.]

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### 222 Role of CD36 in Ozone (O3)-Induced Lung Injury, Inflammation, and Vascular Dysfunction.

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Ground level O3 can damage the cardiovascular system. A lack of a clear mechanism explaining O3-induced vascular health effects hinders the effectiveness of policies for achieving better health. Evidence suggests that inhaled pollutants evoke a systemic inflammatory response that causes endothelial injury and dysfunction. Using serum from O3-exposed mice, we found that circulating components impaired acetylcholine (ACh) vasorelaxation in aortas from naive wild type (WT) mice. However, the mechanistic interaction(s) between circulating factors and endothelial cells is unknown. To address this issue we turned our attention to pattern recognition receptors (PRRs), such as CD36 (cluster of differentiation 36), as mediators of vascular abnormalities following O3 exposure. PRRs are capable of detecting danger signals released by stressed or injured cells. We hypothesized that activation of endothelial CD36 following acute O3 exposure mediates cross-talk between lung-derived circulating factors and vascular endothelium, culminating in endothelial dysfunction.

Female C57 wild type (WT) and CD36 knockout (KO) mice were exposed to filtered air (FA) or 1 ppm O3 for 4 h. Indices of pulmonary (quantified by lavage inflammatory cells) inflammation was assessed 24 h later. The effects of exposure on ACh-induced vasorelaxation were studied using the aortic ring preparation. Parallel experiments were performed in aortas from naive WT mice incubated with serum from exposed mice.

O3-induced infiltration of macrophages and neutrophils into the airspace in WT mice were absent in CD36 KO mice. ACh-evoked vasorelaxation of thoracic aorta of WT mice, but not CD36 KO mice, was significantly reduced after inhalation of O3. Ex vivo assays utilizing homologous serum demonstrated that the vascular damage caused by O3-induced circulating factors was dependent on vascular CD36 receptor expression.

Collectively, our data demonstrate that an as yet unidentified circulating factor, or factors, induced by O3 exposure leads to vascular dysfunction mediated, in part, by CD36 binding in the vascular tissue.
**Interaction of Human Bronchial Epithelial Cells and Alveolar Macrophages Modifies the Innate Immune Response to Ozone.**

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The lining of the airways consists of airway epithelial cells and resident immune cells, which together coordinate the innate immune response to oxidant pollutants. Oxidant pollutants can damage airway epithelial cells and induce the production of soluble mediators that attract nearby immune cells, such as alveolar macrophages, and activate innate immune pathways. Using ozone (O3) as a model oxidant pollutant, we developed a human bronchial epithelial cell (HBE) and alveolar macrophage (AM) co-culture model to assess how the interaction between HBE and AM modifies the innate immune response to oxidant air pollutants. AM derived from the bronchoalveolar lavage of healthy volunteers were co-cultured with the HBE cell line 16HBEo- on transwell cell culture inserts or on transwells alone. Co-cultures, AM alone, and HBE alone were exposed to 0.4 ppm O3 or clean air for 4 hours and analyzed 1 and 24 hours after O3 exposure. O3-induced secretion of interleukin (IL)-1β and IL-8 was compared between the cultures to determine the specific cellular sources and whether the interaction between AM and HBE modifies the inflammatory response. Using flow cytometry, co-cultures or AM alone were examined for AM surface receptor expression, particularly CD44, Toll-like Receptor 4 (TLR4) and its co-receptor CD14, which recognize soluble mediators produced in response to oxidative damage. Our results suggest that co-culture of AM and HBE modifies O3-induced secretion of IL-1β, but not IL-8. Whereas the co-cultures had robust O3-induced IL-1β production, this response was blunted in both AM and HBE cultured alone. Both O3-exposed AM had altered CD14, TLR4, and CD44 expression, and co-culture further modified surface marker expression. These results suggest that HBE and AM coordinate the inflammatory response to O3, and that the interaction between HBE and AM is an important determinant of the innate immune response to inhaled oxidant pollutants.

**Behaviour of Lactose Blends in Nonclinical Respiratory Safety Assessment Studies.**

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Drug/lactose powder blend range and complexity has also grown proportionally as the potency of drug molecules has increased. This review, therefore, addresses the relationship between the active drug moieties and total particulate over a range of lactose powder blend formulations before and after aerosolisation to ensure enhanced study control. Test atmosphere concentration data from studies were chemical or gravimetric analysed for drug:total particulate (TP) ratios and compared with the original blend strength (BS). Results for unmicronised blends indicated up to a 13-fold increase in drug:TP ratio for lower drug blend strengths (0.25% w/w) for non-rodent exposure systems and 9-fold for rodent systems. The fold change (FC) decreased exponentially with increasing drug BS for unmicronised blends and by 40% w/w; the FC was only 2-fold. Greater variation between different drug actives was also evident at lower blend strengths. Using stabilisers or additives resulted in a slight increase (between 0.25 and 0.50 w/w) in the drug:TP ratio over the same BS range. Comparison of 3 different lactose types indicated varying proportions of FC depending on the particle size of the lactose. Micronised Lactohale LH301 gave marginal increase in FC between BS of 0.25 and 40% w/w (1.02 to 1.11) but the FC exhibited for unmicronised Respitose SV008 gave a much greater change over the same BS range. The FC using the same BS with different size inhalation chambers showed no difference with chamber level implying that the animals receive the same drug:lactose ratio irrespective of their position within the chamber. Comparison between the Flow-Through and Flow past chambers with different blend strengths gave a greater FC for the latter chamber type (up to 67%) due to increased lactose sedimentation related to internal geometry of the chamber. In conclusion, this approach will allow greater prediction and confidence that the gravimetric aerosol concentrations will be an accurate representation of the active drug moiety when the frequency of chemical analysis is reduced.

**An In Vitro Cell-Based Assay to Measure the Solubilization of Indium-Containing Particles by Macrophages.**


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Inhaled or airway-delivered indium-containing particles (ICPs) such as indium phosphate (InP) and indium tin oxide (ITO) exhibit pulmonary toxicity and are carcinogenic. Many ICPs are highly insoluble compounds which are engulfed by alveolar macrophages; however, the mechanism(s) of ICP-induced pathogenesis within the lungs is unclear. We have previously shown that ICPs are cytotoxic to macrophages in vitro which is dependent upon phagolysosome acidification. In the current study, we hypothesized that macrophages phagocytose and solubilize ICPs which generates free indium ions—likely the cytotoxic constituent of ICPs. Adherent RAW 264.7 macrophages were treated with 200 μg/ml InP particles for 24 hrs. In some groups, macrophages were pre-treated for 30 min with 5 μg/ml cytoketatin D (cytoD), an inhibitor of phagocytosis, and then co-treated with InP + cytoD for 24 hrs. CytoD treatment blocked both the phagocytic uptake and cytotoxicity of InP particles. Cell culture supernatants were collected after 24 hrs of treatment and centrifuged to pellet any residual cells and InP particles. Cell and particle-free supernatants were then acid digested overnight and the concentration of total indium was measured using atomic absorption spectroscopy. The concentration of extracellular indium in cell culture supernatants from macrophages treated with InP particles in the absence of cytoD (91.2 μg/L) was significantly increased and approximately 3-fold greater compared to macrophages treated with InP in the presence of cytoD (29.8 μg/L). These data indicate that macrophages phagocytose and solubilize ICPs within lysosomes resulting in cell death and the extracellular release of free indium metal species. This cell-based assay can potentially be applied to other ICPs to determine if the in vitro macrophage solubility of different ICPs correlates with in vivo pulmonary toxicity.

**Acute Pulmonary Heme Oxygenase-1 Protein Response to Particulate Matter in Naïve Animals As an Indicator of Allergenic Adjuvant Potential.**


The San Joaquin Valley (SJV) of California is home to high PM pollution and asthma symptom prevalence. Recently, source oriented sampling (SOS) approaches have been employed in the SJV to collect commonly occurring particle source combinations in the normal milieu of PM typical to the region to allow for the evaluation of their differential toxicities. Acute toxicity studies to assess differential pulmonary inflammation were performed. Additional studies were performed in an acute model of allergic airway inflammation. Studies utilized BALB/c mice, 8-10 weeks old and re-suspended particles collected and extracted by SOS methods. Naive studies utilized dosing via oropharyngeal aspiration of 50 μg SOS PM with tissues examined for pulmonary inflammation 24-hours post dosing. Allergic studies utilized intranasal aspiration dosing on day 1, 5, and 5 g of a) vehicle control, b) 25μg endotoxin purified D. Farinae house dust allergen, HDM (allergic control), or c) HDM and 15μg SOS PM (45μg total dose). Animals were challenged on day 11, 12, and 13 with allergen alone and tissues collected on day 14. All allergen-treated animals exhibited cellular profiles indicative of an allergic response with elevations in leukocytes characterized by neutrophils, lymphocytes and eosinophils. Heme oxygenase-1 (HO-1) protein levels in homogenized pulmonary tissue of acutely exposed naïve mice were quantified as a biomarker of oxidative stress. Analysis of correlations revealed large associations between total cell, lymphocytic, eosinophilic and neutrophilic pulmonary inflammation in allergic animals in contrast to HO-1 protein levels in the tissue of acutely exposed naïve animals. Cellular inflammation in naïve acute studies did not correlate with HO-1 protein and did not accurately predict adjuvant potential. These studies suggest that pulmonary HO-1 levels in naïve acute studies and existing archived tissue may be a valuable indicator of particle adjuvant potential. Support: CARB/EPRI

**Toxic and Mutagenic Effects of World Trade Center Dust on Cultured Human Lung Cells.**

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The terrible events of the September 11, 2001 World Trade Center (WTC) tragedy left many dead, injured, devastated and emotionally scarred. The effect of this tragic event was noticed even a decade later in many upper-respiratory complica-
Dose Response to Sacramento Particulate Matter.


Particulate matter (PM) exposure contributes to respiratory diseases and cardiopulmonary mortality. The toxicity of PM could be related to sources, such as vehicle exhaust, and composition, such as abundance of polycyclic aromatic hydrocarbons (PAHs). We exposed adult male balb/c mice, via oropharyngeal aspiration, to a range of doses of PM2.5 collected during the winter in downtown Sacramento near a major freeway interchange (SacPM). Because the relative contribution of PAHs in SacPM might be important, and since filter extraction may alter PM biological effects, we tested two PM preparation methods (sonication/spin-down and sonication/lyophilization) at 10, 50 and 100 μg doses and analyzed the lung tissue response at 24 hrs after dosing. We analyzed 1) leukocytes and total protein in BALF, 2) airway-specific and whole lobe expression of PAH sensitive genes (CYP1B1 and CYP1A1) and IL-1b and 3) lung histology. We found both PM extraction methods stimulated similar biological responses, but the spin-down method was more robust at producing IL-1b and CYP1B1 gene responses and the lyophilization method induced whole lung CYP1A1. Neutrophils in the BALF were increased 5-fold at 50 μg and 10-fold at 100 μg. Total protein in the BALF was significantly increased at both the 50 and 100 μg doses. Histopathology scores were dose responsive and more robust in mice treated with spin-down derived PM. CYP1B1 gene expression in whole lung increased 3-fold at the 50 and 100 μg dose for this method as well, but was increased less than 1.5-fold for the lyophilization method. In microdissected airways all dosages of the spin-down PM increased CYP1B1 gene expression significantly, but the lyophilized PM did not change CYP1B1. We conclude 1) the method of filter extraction can influence the degree of biological response at a given dose, 2) for SacPM the minimal effective dose for this strain of mouse and route of exposure is 50μg doses. Histopathology, score, and lung parenchyma have differential ability to be upregulated in response to PAH-containing PM. (Supported by California Air Resources Board)

230 Subchronic Inhalation Exposure of Rats to Libby Amphibole and Asbestos: Effects at 1 and 3 Months Postexposure.

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Increased asbestosis, lung cancer, and mesothelioma rates are evident after exposure to Libby amphibole (LA). To support dosimetry model development and compare potency, a subchronic nose-only inhalation study (6 hr/d, 5 d/wk, 13 wk) was conducted in male F344 rats. Rats were exposed to air (control), LA (LO, MED, HI; 1:01, 3:33, 10.08 mg/m3; 300, 701, 4050 fibers/cc), or asbestosis (AM; 3.35 mg/m3; 1035 f/cc). Toxicity endpoints, pathology, and fiber burden evaluation were determined 1 d and 1, 3, and 18 mo post-exposure. Previously reported results (Dodd, SOT 2013) showed comparable inflammatory and fibrogenic responses 1 d after exposure to MED LA and AM. Here we report BAL neutrophils were increased in HI LA and AM groups at 1 mo, but only in HI and MED LA groups at 3 mo. Macrophages were decreased in the HI and MED LA groups at 3 mo. AM and MED LA groups had comparable increases in BAL LDH and protein at 1 and 3 mo. Lung tissue cytoplasmic production and activation of pro-fibrotic and cellular growth signaling pathways were also comparable between AM and MED LA. Histopathological examination of the lung found a minimal increase in alveolus inflammation, interstitial fibrosis, bronchiolization, and foreign body presence in both AM and LA groups compared to controls. Alveolus inflammation was most severe in HI LA rats at 1 and 3 mo compared to all other exposure groups. Only HI LA rats had bronchiolitis epithelial hyperplasia, but only 1 mo after exposure. These results show comparable fibrogenic responses 1 and 3 mo after subchronic exposure of rats to LA and AM asbestos. Tissue fiber burdens are being measured to support dosimetry model development of deposition and clearance (Ashgarh, SOT 2012: Jarabek, SOT 2013); comparison of responses between fibers may change based on dosimetry modeling. (This abstract does not represent US EPA policy)

231 Exposure to Sumas Mountain Chrysotile Induces Similar Gene Expression Changes As Libby Amphibole but Has Greater Effect on Long-Term Pathology and Lung Function.

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This study was designed to provide understanding of the toxicity of naturally occurring asbestos (NOA) including Libby amphibole (LA), Sumas Mountain chrysotile (SM), El Dorado Hills tremolite (ED) and Ontario ferroactinolite cleavage fragments (ON). Rat-respirable fractions (aerodynamic diameter ≤ 2.5 μm) were prepared by water elutriation and a dose of 1.5 mg/rat delivered via a single intratracheal (IT) instillation. Bronchoalveolar lavage (BAL), gene expression, histopathology, lung function and histopathology were analyzed 1 d, 3 mo, or 15 mo post-instillation.

One day after exposure, although inducing less acute inflammation than other samples, LA and SM induced a greater degree of lung injury. A similar trend was also observed in gene expression profiles, as both LA- and SM-exposed rats differed significantly from dispersion media (DM) controls. Changes were suggestive of dysregulation of both extracellular matrix and fibrosis pathways. By three months, most BAL parameters had returned to DM control levels. However, significant time-dependent fibrosis was evident in rats exposed to LA or SM. By 15 months, the greatest fibrotic changes were observed in SM-exposed rats; while no fibrosis was noted in the cleavage fragment or DM control group (SM>LA>ED>ON=DM). Consistent with the greatest degree of fibrosis, only SM-exposed rats exhibited persistent, long-term changes in lung function parameters. These data demonstrate that, in the rat, SM resulted in more significant long-term effects after a single IT exposure than LA. This study suggests that there may be cause for concern for people at risk of being exposed to NOA from the Sumas Mountain landslide, and highlights the need for further study of sites where NOA is present. (This abstract does not represent U.S. EPA policy)

232 Lack of Sex Difference in Nasal Glutathione Response to Naphthalene.

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At concentrations of 10 ppm or greater, naphthalene (NA) is a nasal carcinogen inducing respiratory adenomas in male and olfactory neuroblastosomas in female rats, respectively. The proposed carcinogenic mode of action includes metabolic activation via CYP450 to an electrophile with subsequent ethane-ethylene-depiction, escape of electrophile, and covalent binding. Respiratory and olfactory mucosa are tumor target sites and both contain NA CYP450 activating capacity; the activity in olfactory may exceed that in respiratory mucosa by 3-fold. To fully define the effect of NA on nasal glutathione, male and female F344 rats were exposed to 0, 10, or 30 ppm NA for 1, 4, or 6 hours. Following exposure, nasal olfactory and respiratory tissues were analyzed for reduced/oxidized glutathione levels (GSH/GSSG). Female control rats had twice the levels of GSH compared to male controls, but NA exerted similar effects on GSH in both genders. GSH was depleted at all times; in females,
respiratory and olfactory mucosal levels were ~45 and 70% of control levels (re-
spectively) after 10 or 30 ppm, an effect that did not correlate with local CYP450 
activation rates. Similar trends were seen for male rats. To fully define the concen-
tration, rats were exposed to 0, 0.5, 1, 3, 10, or 30 ppm NA for 4 hours. 
Significant GSH depletion occurred at all exposure levels in respiratory and olfac-
tory mucosa with maximal depletion (about 30 and 60% of control levels, respec-
tively) occurring at or above 1 ppm. Similar trends were seen in both male and 
female NA, at concentrations well below those shown to be carcinogenic, causes 
significant depletion of GSH in the nose. The degree of GSH depletion in different 
nasal regions does not correlate with activation rates, suggesting that other factors 
contribute to the GSH response. No sex difference was observed in GSH response, 
suggesting that the sex difference in tumor response cannot be attributed to this 
step in the carcinogenic mode of action. (Supported by the Naphthalene Research 
Council)

234 Upper Respiratory Lesions in Rats Administered 
Amiodarone Hydrochloride Solution Orally for 4 Days by 
Intrasacofugal Dosing: Absence of the Lesion by 
Intragastric Dosing.

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Upper respiratory (UR) tract can be damaged by compounds administered orally. 
Retrospective exposure of nasal passage to dose formulation from the esophagus has 
been suggested as one of the toxicological mechanisms (Damsch et al. Toxicol 
Pathol, 2011). However, literature are limited with this toxicity of orally adminis-
tered drugs in clinical use and the toxicological significance remains unclear. 
Amiodarone hydrochloride (AM) is an antiarrhythmic agent administered orally 
and intravenously in clinical use. We demonstrated AM induces UR lesions in rats 
by gavage dosing for 4 days with a metallic tube. Retrospective exposure of the nasal 
passage to dose formulation was suggested because incidence of the lesion was 
higher in the posterior nasal passage than the anterior one (Ogata et al., JSTP 
Annual Meeting, 2011). Furthermore, irritability of the dose formulation to nasal 
epithelium was confirmed based on induction of UR lesions after single intranasal 
dosing (Ogata et al., JSOT Annual Meeting, 2012). To examine the effect of the 
doing procedure on the lesions, rats were administered 150 mg/kg AM by a 
catheter instead of a metallic tube and the UR lesions were compared histopatho-
logically with those obtained in the former study with a metallic tube. In this study, 
no UR lesions were observed in rats given AM with a catheter. Cmax and AUC was 
equivalent between dosing procedures with the catheter and metallic tube after a 
single dose. The results suggest that the UR lesions in rats given AM by a metallic 
tube is not attributable to systemic exposure alone. Dependence on metallic tube 
dosing may suggest low toxicological relevance of the UR lesions. Furthermore, 
catheter dosing would be an option to evaluate toxicological significance of gavage-
related UR lesion, with equivalent Cmax and AUC obtained by metallic tube dos-
ing, and without possible concern specific to dosing from parenteral or feeding 
routes.
Diacetyl-Induced Respiratory and Olfactory Toxicity in Mice: Influence of Ubiquitination, Gender, and Dichacarbonyl/L-Yxylolox Reductase Gene Knockout.

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The ét-dicarbonyl butter flavoring, diacetyl (2,3-butanedione), is associated with flavorings-related constrictive bronchiolitis in workers who make or use flavorings. Diacetyl causes protein damage in a process believed to be dependent upon the ét-dicarbonyl structure. A protective response to damaged protein is ubiquitination with subsequent proteasomal processing. Diacetyl is also metabolized to the less reactive ét-hydroxyacetone, acetoin, by dicarbonyl/L-yxylolox reductase (Dctx). We examined the role of Dctx and gender on acute toxicity of inhaled diacetyl by exposing Dctx knockout and wildtype mice of both sexes to diacetyl at target concentrations of 0, 100, 200 or 300 ppm for 6 hr. At 1 day post-exposure, endpoints were semi-quantitative histopathology and morphometric measurement of ubiquitin immunohistochemistry in nose and lung sections. Ubiquitin was principally localized to nasol and intrapulmonary airways, increased in large bronchioles at concentrations ≥100 ppm, and in the nose at 300 ppm. Diacetyl-induced ubiquitin in the nose and lung was modified by both gender and Dctx. In lung histopathology, diacetyl caused vacuolation of airway epithelium of large bronchioles at concentrations ≥100 ppm. In olfactory bulb (OB) of male mice inhaling 300 ppm diacetyl, mRNA expression of inflammatory mediators and olfactory marker protein (Omp), a marker of olfactory neuron axons, were assayed by real-time PCR. Diacetyl elevated Il6, Cxcl2, and Tnfa and decreased Omp in OB. The data suggest that ubiquitin expression is a sensitive biomarker of diacetyl-induced protein damage in airway epithelium. Further, diacetyl causes neuroinflammation and potential loss of axons of olfactory neurons in OB, suggestive of neurotoxicity.

Subchronic Exposure to Ambient Particulate Matter Induces Oxidative Stress Responses in Brain Tissue of ApoE-/- Mice.

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Exposure to particulate matter (PM), present in urban environments, has been shown to induct pro-inflammatory and oxidative stress responses in the central nervous system (CNS) of apolipoprotein E knockout (ApoE-/-) and Balb/c mice. In this study oxidative stress responses in different subcellular fractions of the ApoE-/- mouse brains were evaluated after a subchronic exposure to fine (≤2.5 μm) concentrated ambient particles (CAPs). Apo E-/- mice were exposed to either CAPs or particle-free air for 5 hours a day, 5 days per week, for a period of 6 months. The whole-body inhalation exposures were conducted in two urban cities (Seattle, WA and Detroit, MI) with distinct sources and chemical composition of PM. Brain tissue was collected after the exposures were completed and analyzed for biomarkers of oxidative stress. The antioxidant glutathione was reduced in the brains of mice exposed to UFP+O3 compared with exposure to either PM2.5 or UFP alone. This suggests that the in vivo exposure to UFP+O3 would enhance the acute effects of particles, particularly UFP. Conscious unrestrained C57BL/6 mice implanted with radiotelemeters were exposed by whole-body inhalation to either 250 μg/m3 PM2.5 or 100 μg/m3 UFP with or without 0.3 ppm O3 (4hrs); separate groups were exposed to either filtered air or O3 only. Heart rate (HR) and electrocardiogram (ECG) were recorded continuously before, during and after exposure. Control animals experienced a decrease in HR during exposure. Neither PM2.5 or UFP alone caused any HR change; however with O3 co-exposure, HR remained transiently elevated above control levels. Exposure to UFP+O3 caused decreased PR-interval, a transient increase in QRS, and increased QTc. PM2.5 alone caused QRS to decrease and O3 alone caused a decrease in QRS interval and QTc. There were no other significant differences in the ECG parameters measured of any groups. Lastly, only animals exposed to UFP+O3 had an increase in the number of non-conductive P-waves; there were no differences in other arrhythmia counts. These data suggest that O3 co-exposure might worsen the stress response to PM, especially UFP, and cause repolarization heterogeneity in the heart, which increases the risk for arrhythmogenesis. As such, this indicates that the cardiovascular effects of particle and gas co-exposures are not easily characterized, potentially increasing the complexity of risk assessment. (This abstract does not reflect EPA policy).

Co-Exposure to Ultrafine Particulate Matter and Ozone Causes Electrocardiogram Changes Indicative of Increased Arhythmia Risk in Mice.

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Numerous studies have shown a relationship between acute air pollution exposure and increased risk for cardiovascular morbidity and mortality. Due to the inherent complexity of air pollution, recent studies have focused on co-exposures to better understand product-4-hydroxyalkenal (HNE). This study was designed to evaluate the cardiac effects of concentrated ambient fine (PM2.5) and ultrafine (UFP) particles with and without ozone (O3) co-exposure. We hypothesized that ozone co-exposure would enhance the acute effects of particles, particularly UFP. Conscious unrestrained C57BL/6 mice implanted with radiotelemeters were exposed by whole-body inhalation to either 250 μg/m3 PM2.5 or 100 μg/m3 UFP with or without 0.3 ppm O3 (4hrs); separate groups were exposed to either filtered air or O3 only. Heart rate (HR) and electrocardiogram (ECG) were recorded continuously before, during and after exposure. Control animals experienced a decrease in HR during exposure. Neither PM2.5 or UFP alone caused any HR change; however with O3 co-exposure, HR remained transiently elevated above control levels. Exposure to UFP+O3 caused decreased PR-interval, a transient increase in QRS, and increased QTc. PM2.5 alone caused QRS to decrease and O3 alone caused a decrease in QRS interval and QTc. There were no other significant differences in the ECG parameters measured of any groups. Lastly, only animals exposed to UFP+O3 had an increase in the number of non-conductive P-waves; there were no differences in other arrhythmia counts. These data suggest that O3 co-exposure might worsen the stress response to PM, especially UFP, and cause repolarization heterogeneity in the heart, which increases the risk for arrhythmogenesis. As such, this indicates that the cardiovascular effects of particle and gas co-exposures are not easily characterized, potentially increasing the complexity of risk assessment. (This abstract does not reflect EPA policy).


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Classically in vitro diesel exhaust (DE) studies have been performed with submerged cell cultures which are exposed to collected DE particles. Major drawback of this exposure method is that cells are not exposed to the whole complex and dynamic mixture of compounds DE is composed of. In recent years, air-liquid interface exposures have become more widely used, enabling in vitro exposures to mixtures of gases and particles. The main objective of this study was to investigate the feasibility of exposing human lung epithelial cells at the air-liquid interface to complete DE generated by a heavy-duty truck in the state-of-the-art TNO power train facilities. Human epithelial lung cells (A549) were directly exposed at the air liquid interface to DE generated by a heavy-duty Euro III truck (turbo diesel model 2002). The truck was tested at a steady-state cycle at a speed of ~70 km/h-1 to simulate free-flowing traffic at a motorway on a transient engine dynamometer. Cells were exposed to DE for 1.5 hours. After a 24 hours post-incubation period, cells were analysed for markers of oxidative stress (glutathione levels, GSH; heme oxygenase 1 protein levels, HO-1), cytokotoxicity (lactate dehydrogenase release, LDH; Alamar Blue assay) and inflammation (interleukine-8 protein levels, IL-8). DE exposure resulted in a decreased cell viability (significantly decreased Alamar Blue levels in the incubation medium and slightly increased LDH levels), and an increased oxidative stress response (significantly increased HO-1 levels and reduced GSH/GSSG ratio). However, the pro-inflammatory response seemed to decrease (non-significant decrease in IL-8).

The presented results here demonstrate that our in vitro exposure approach is indeed well suited for testing complex particulate and gaseous pollutant mixtures from diesel trucks. Our results confirm previous in vitro studies showing cytotoxicity and oxidative stress responses due to DE exposure.

Comparative Toxicity of Soy Biodiesel and Diesel Emissions in Healthy and Allergic Mice.

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Toxicity from combustion of 100% soy-based biodiesel (B100) was compared to that of petro diesel (B0) or a 20% biodiesel / 80% petro diesel mix (B20) in healthy and house dust mite (HDM)-allergic Balb/c mice. Exhaust from combustion of B0, B20 and B100 was diluted with 50% or 500 μg/m3 as determined by real-time Tapered Element Oscillating Microbalance. Studies in healthy mice showed greater levels of MIP-2 and neutrophils in bronchoalveolar lavage (BAL) fluid 2 hr after a single 4 hr exposure to B0 compared with exposure to biodiesel emissions (air control neutrophils = 1×, B0 = 11.9×, B20 = 4.4×, B100 = 2.1×). However these differences were attenuated 24 hr after exposure and no consistent differences were observed 2 or 24 hr after 5 d (4 h/d) or 4 wk (5 d/wk) exposures. Mice sensitized and challenged intranasally with HDM and exposed to B0, B20, or B100 for 4 wk (5 d/wk) had no emissions-related differences in airway...
hyperresponsiveness to MCh aerosol (determined by total lung resistance in anesthetized, paralyzed and ventilated mice). Non-significant trends of decreased eosinophils and IL-5 in BAL fluid were found after exposure to the 500 µg/m3 concentration of all 3 fuels. Proliferative responses of peribronchiolar lymph node cells in response to HDM antigens in vitro were not significantly affected by exposure to fuel emissions. We conclude that alternative soy biofuel emissions have comparable or reduced adverse effects relative to diesel emissions in healthy mice or a mouse model of allergic asthma. (This abstract does not represent U.S. EPA policy.)

### Mechanisms Underlying Anti-Inflammatory Effects of Selective Diindolylmethane Compounds Using RAW264.7 Cells.

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Chronic inflammation has been associated as the root cause of many serious illnesses including heart disease, Alzheimer’s disease and many cancers. Tolfenamic Acid (TA) a non-steroidal anti-inflammatory (NSAID) drug has been shown to have multiple anti-inflammatory effects in RAW264.7 murine macrophages. This study compares the anti-inflammatory effects of selected diindolylmethane (DIM) compounds in activated RAW264.7 cells. Results indicate a significant decrease in production of prostaglandin E (2), D (2) and F (2). In addition, there was a decrease expression of mediators of inflammation including cyclooxygenase-2 (COX-2) in LPS-induced RAW 264.7 cells. These results underscore the potential use for these DM compounds in ameliorating inflammation in disease processes and therapeutic regimens.

### Fyn Kinase Inhibitors Attenuate Manganese Nanoparticles Induced Neuroinflammatory Signaling in BV2 Microglial Cells and Primary Microglia.

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Manganese (Mn) nanoparticles are currently used in a multitude of industrial and biomedical applications, including magnetic resonance imaging, high capacity batteries, industrial coatings, biosensors, plastics, ultrahigh density storage devices, nanofibers, and catalysts. Yet, the potential impacts of these particles on human health and the environment are not well understood. Notably, the cellular mechanisms underlying Mn nanoparticle induced neurotoxicity are yet to be identified. Because nanoparticles are similar in size to microbes, we hypothesize that Mn nanoparticles exposure may activate phagocytic microglial cells to induce a neurotoxic response. We exposed BV2 microglia and primary microglial cultures to various doses of Mn nanoparticles and then measured inflammatory markers. Exposure of 0-50 µg/mL Mn nanoparticles to BV2 microglia over 24 hr period resulted in a dose-dependent increase in NO, ROS, TNFβ, IL-6, IL-12, and RANTES levels. Additionally, Mn nanoparticles induced over a threefold increase in ROS generation in primary microglial cells. Interestingly, Mn nanoparticles also activated the non-receptor tyrosine kinase Fyn in BV2 microglia. In order to determine whether Fyn kinase plays a role in the Mn induced inflammatory response, we tested a series of Fyn kinase inhibitors, including rosinic acid, dinitorosinic acid and caffeic acids. Rosmarinic and caffeic acids showed EC50s of 33 and 32 µM, respectively, against Mn-induced iNOS activation. In addition, Fyn kinase inhibitors significantly blocked Mn-nanoparticle induced TNFβ, IL-6, IL-12 release and ROS generation, indicating that Fyn kinase may play a central role in the Mn nanoparticle neuroinflammation and oxidative stress. Taken together, our results demonstrate that Mn nanoparticles activate microglial cells via a Fyn kinase dependent mechanism and that Fyn kinase inhibitors may serve as efficacious therapeutic agents against the metal nanoparticle induced neuroinflammatory insult (supported by NH grants ES10586 and NS65167).

### S-Adenosylhomocysteine Inhibits NF-κB Activity in the Nucleus of Hepatocytes and Confers Sensitivity to TNF Cytotoxicity.

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Sepsis is a syndrome of infection complicated by vital organ dysfunction and considered one of the leading causes of mortality in intensive care units. It is characterized by a generalized inflammatory response caused by systemic activation of macrophages and other immune and microvascular endothelial cells. The present study demonstrates the efficacy of a novel anti-inflammatory peptide termed IIIIM1 in ameliorating sepsis induced by the bacterial endotoxin lipopolysaccharide (LPS). A single injection of the peptide 3 days prior to lethal dose of LPS increased animal survival by 60%. Shorter or longer intervals between IIIIM1 and LPS administration reduced peptide efficacy. LPS-induced increase in liver, kidney and spleen mass was reverted by peptide treatment. Thioglycolate-induced peritonitis, expressed by increased number of macrophages in the peritoneal cavity, was reduced by 38% in IIIIM1-treated mice. A bell-shaped dose-response effect was observed in both peritonitis and sepsis models reaching maximal effect at 1mg/kg. Chemical modifications such as omission of the C- or N-terminal residues weakened the anti-inflammatory activities of the peptide. Serum levels of interleukin 6 and tumor necrosis factor alpha were reduced by 32% and 53%, respectively, in IIIIM1 treated mice intoxicated by LPS. Similar cytokine profile was not observed in LPS-activated peritoneal macrophages treated in vitro with the peptide. In view of these data IIIIM1 is a promising drug candidate for treatment of sepsis induced by bacterial endotoxin.

### Proinflammatory Cytokines Present in the Tumour Microenvironment Induce Phenotypic Change in Colorectal Cancer Cell Lines.

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Colorectal cancer (CRC) is the third most common cancer worldwide with metastatic disease responsible for high mortality rates. The cellular microenvironment is modified during malignancy to support tumor development and metastasis, however mechanisms by which this occurs remain unclear. Previous data from our laboratory has shown an overexpression of cytochrome P450 (CYP) drug metabolizing enzymes in CRC cells treated with thiophenolone (THP1) and tumor necrosis factor alpha (TNFα) and IL-6 in neoplastic tissue resected from colorectal cancer patients compared to matched non-neoplastic controls. In the current study, the presence of these cytokines in the tumour microenvironment was investigated in vitro using two CRC cell lines HCT116 and SW480, differing in metastatic potential. Treatment with conditioned media from activated THP1 monocytes, known to secrete an array of cytokines (including IL6, IL1β and TNFβ), was able to induce the invasive properties of the metastatic cell line HCT116 as measured using wound and migration chamber assays. Treatment with IL6 alone (0-1000 pg/ml) was also shown to promote cell motility and invasion in a bell-shaped dose response, characteristic of cytokine functionality. The non-metastatic cell line SW480 did not respond to the conditioned media and was more resistant to IL6, generating a response only at the higher doses. Additionally, CYP 1B1 and 2E1 expression were increased following treatment with the conditioned media as well as with IL6 on its own. Taken together, these data indicate that pro-inflammatory cytokines, in particular IL6, are able to cause phenotypic change in CRC cells by inducing CYP expression and promoting ability for the cells to migrate and invade surrounding tissue, thus demonstrating the important role of tumor microenvironment in disease progression.

### A Novel Anti-Inflammatory Peptide: Potential Therapy of Sepsis Induced by Bac terial Endotoxin.

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Chronic alcohol exposure results in liver injury that is largely driven by inflammatory cytokines such as tumor necrosis factor-alpha (TNF). Hepatocytes are normally resistant to the cytotoxic effects of TNF, but they become sensitized to TNF by chronic alcohol exposure. Recently we reported that the decrease in the ratio of S-adenosylmethionine (SAM) to S-adenosylhomocysteine (SAH) that occurs with alcohlc liver injury renders hepatocytes sensitive to TNF cytotoxicity. The purpose of the present study was to determine whether inhibition of the transcription factor NF-kB contributed to TNF-induced cell death in hepatocytes with high levels of SAH. Hepatocytes were pre-incubated with LPS 3 days prior to lethal dose of LPS and then incubated with IIIM1 in an attempt to decrease SAH levels and then incubated with IIIM1 in an attempt to decrease SAH levels. Hepatocytes were then incubated with LPS for 24 hr. Following stimulation with TNF, viability was determined by the MTT assay, and activation of the NF-kB pathway was assessed by measuring degradation of cytosolic IkBα, translocation of NF-kB to the nucleus, and expression of an NF-kB-dependent reporter gene as well as endogenous targets of NF-kB related to cell death. The results showed that NF-kB was indeed inhibited in cells with high SAH, and that this inhibition occurred at the level of the nucleus; the cytoplasmic inhibitor IkBα was degraded and NF-kB translocated to the nucleus in response to TNF stimulation of HepG2 cells in both control cells and in...
cells with high SAH levels. Nuclear NF-kB was not transcriptionally active, how-
ever, when SAH levels were high. As a result, iκB-α was not re-synthesized and NF-
kB remained in the nucleus. It is likely that cross-talk with other transcription fac-
tors is perturbed under these conditions, resulting in still other changes in gene
expression.

247 Cyanobacterium Anabaena sp. Lipopolysaccharide (LPS) Elicits Release of MIP-α1, Interleukin-6, and Matrix Metalloproteinase-9 from Rat Brain Microglia In Vitro.

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We recently reported that freshwater cyanobacterium Anabaena (AnaLPS) elicited release of superoxide anion (O2·-), thromboxane B2 (TXB2), and tumor necrosis alpha (TNF-α) by rat microglia (BMG) in vitro (The Toxicologist CD 126 (S-1), 2012). We hypothesized that AnaLPS-activated BMG might additionally re-
lease the cytokine interleukin-6 (IL-6), the chemokine MIP-1α (MIP-1α), and the matrix metalloproteinase-9 (MMP-9) in vitro. Methods: AnaLPS was prepared by hot phenol/water extraction. BMG were isolated from neonatal rats, and treated in vitro with 0.1-105 ng/mL AnaLPS at 35.9 °C for 17 hours. TNF-α, MIP-1α and IL-6 were determined by Milliplex® MAP rat cytokine/chemokine immunoassays, and MMP-9 by zymography. Results: MIP-1α, TNF-α, and IL-6 generation were observed at AnaLPS >10 ng/mL (MIP-1α), and > 1,000 ng/mL, respectively. In contrast, MMP-9 release was significant at AnaLPS 10,000 ng/mL. Conclusions: Treatment of BMG with AnaLPS confirmed previously reported TNF-α release, and furthermore demonstrated for the first time the release of MIP-1α, IL-6 and MMP-9. Taken together, our results suggest that in vitro proinflammatory media-
tor release by cyanobacterial AnaLPS-activated BMG is complex, including liquids (TXB2), free radicals (O2·-), cytokines (TNF-α & IL-6), chemokines (MIP-1α) and enzymes (MMP-9), all of which might play a yet unknown role in the putative im-
munotoxicity of AnaLPS in vivo. Continued investigation of AnaLPS chemistry and immunotoxicology are currently ongoing in our laboratories. Supported by Midwestern University and the University of Hawaii at Manoa.

248 Spleen As a Source of Inflammatory Macrophages: Role in Acetaminophen-Induced Hepatotoxicity.

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Activated macrophages (MP) have been implicated in the hepatotoxicity of acet-
aminophen (APAP). However, the origin of these cells has not been established. Splenic monocytes (mono/MP) have been shown to accumulate at inflammatory sites following tissue injury. In the present studies, we analyzed the contribution of splenic mono/MP to liver inflammation and injury induced by APAP. Mice were fasted overnight prior to administration of APAP (300 mg/kg, i.p.) or PBS. Spleen, bone marrow (BM) and liver were collected 24-96 h later and analyzed by flow cy-
tometry and immunofluorescence for the presence of activated mono/MP. APAP intoxication was associated with a time-dependent increase in CD11b+Ly6C+ proinflammatory MP, and a decrease in F4/80+ resident MP in the liver; this was correlated with a significant decrease in CD11b+F4/80-Ly6C-Ly6G- resident mono/MP in the spleen at 24 and 48 h post APAP, with no effect on BM mono/MP. Conversely, CD11b+F4/80-Ly6G+ inflammatory mono/MP increased in the spleen after 48–96 h, but decreased in the BM. To assess the role of splenic mono/MP in APAP hepatotoxicity, we used splenectomized (spx) mice. APAP-in-
duced hepatotoxicity was significantly decreased in spx mice, as measured by de-
creases in serum transaminases. Histologic evidence of hepatic necrosis was also re-
duced. This was associated with a significant decrease in inflammatory mono/MP subsets (CD11b+F4/80-Ly6G+ and CD11b+F4/80-Ly6G-) in the BM at 24 h post APAP when compared to sham control mice. In addition, in spx mice, a de-
crease in CD11b+F4/80-Ly6G+ myeloid derived suppressor cells (MDSCs) was observed in BM at 96 h post-APAP. Taken together, these results indicate that splenic mono/MP contribute to early inflammation and hepatotoxicity induced by APAP. Moreover, removal of the splenic reservoir of mono/MP results in emigration of MDSCs out of the BM following APAP administration and this may contribute to tissue repair. Supported by NIH GM034310, ES004738, CA132624, AR055073, ES007148, ES009022.

249 Exaggerated Hepatotoxicity of Acetaminophen (APAP) following Administration of Clodronate Liposomes Is Associated with the Persistence of Classically-Activated Macrophages in the Liver.

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Toxic doses of APAP are known to cause centrilobular hepatic necrosis. Evidence suggests that classically and alternatively activated macrophages play distinct roles in APAP-induced hepatotoxicity. In the present studies, we investigated the effects of macrophage depletion using clodronate liposomes (CL) on activated macrophages currently populating the liver in response to APAP intoxication. Mice were administered empty liposomes (EL) or CL (100 pl, i.v.) 48 h be-
fore APAP (300 mg/kg, i.p.) or PBS control. In mice pretreated with EL, increases in serum transaminases and hepatic necrosis were observed within 24 h; this was correlated with hepatic accumulation of CD11b+Ly6G+ classically activated macrophages in the liver and increased expression of galectin-3, a marker for these cells. These effects were significantly increased in mice pretreated with CL at 72 and 96 h post APAP, and were associated with increases in CD11b+Ly6C+ F4/80/in-
flammatory monocytes in the bone marrow, suggesting a potential origin of these cells. In contrast, APAP-induced expression of the alternative macrophage activa-
tion marker Fizz-1 was reduced by CL, but remained in the nucleus after 24 h, suggesting that increased APAP-induced hepatotoxicity following macrophage depletion using CL is due, in part, to persistent accumulation of classically activated macrophages in the liver. Supported by NIH GM034310, ES004738, CA132624, AR055073 and ES005022.

250 Bone Marrow Inflammation Precedes Delayed Myelosuppression from Hexahydro-1-Nitroso-3, 5-Dinitro-1, 3, 5-Triazine (MNX) Induced in Rats.

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MNX (hexahydro-1-nitroso-3, 5-dinitro-1, 3, 5-triazine) is an environmental ni-
trored product of munitions RDX, contaminates military sites. Our previous stud-
ies identified bone marrow (BM) and spleen as hematological targets of acute
toxicity to MNX in rats in the form of splenic hemosiderosis and loss of BM
Granulocyte Macrophage Colony Forming Cells (GM-CFCs). To address whether delayed loss of GM-CFCs and blood granulocytes (NOAELs 24, 47 mg/kg resp) at 14 days after exposure to MNX is due to persistence of early hematological effects or is late-onset due to required expression period, female Sprague-Dawley rats were orally gavaged with MNX from 0 to 94 mg/kg and different toxicological endpoints were evaluated over a time course at 2, 7, 10, 12, and 14d. Significant de-
crease in relative spleen weight and increased macrophage activity in splenic red pulp at 2d (≥ 47 mg/kg); and persistent splenic hemosiderosis at 2d (≥ 47 mg/kg), 7d (NOAEL 24 mg/kg) and 14d were observed. A significant increase in blood granulocytes and circulating levels of RANTES, a leukocyte chemokine, indicate that an acute inflammatory response occurs at 2d after exposure to MNX. Also, persistent BM macrophage infiltration was observed in MNX (94 mg/kg) treated rat iliums (24h, 2d and 10d) and activation of NFKB signaling pathway in BM cells was evident at 10d. Further, significant increase in adherent BM mesenchymal stromal cell colony forming macrophages, endothelial cells and fibroblasts was ob-
served in MNX (94 mg/kg) treated rats. Collectively, these data suggest that while splenic effects are early onset and persist for at least 14d, myelosuppression is de-
layed until 10d presumably due to development of inhibitory effects of preceding BM inflammation on myelopoiesis. (Support: DoD/CDMRP, US Army Corps of Engineers)

251 Reversibility of Prostate Fibrosis in Response to Bacterial-Induced Chronic Inflammation.

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Introduction: Benign prostatic hyperplasia (BPH) and its associated lower urinary tract symptoms (LUTS) are common in aging men. Chronic inflammation and in-
creased stromal collagen are common features observed in BPH. Recent study has
shown that the collagen content is correlated with the degree of LUTS and tissues status, suggesting that prostate fibrosis in BPH/LUTS. Using the bacterial-prostatic inflammation mouse model, we have shown that prostatic inflammation induces collagen content. The goal of this study is to investigate the reversibility of prostate fibrosis during bacterial-induced chronic inflammation.

Methods: We transurethrally instilled uropathogenic E-coli into adult C3H/HeOuJ male mice to induce chronic inflammation in the prostate. Naïve, saline and E-coli instilled animals were sacrificed after 1 month. Other animals were treated with Baytril in drinking water to resolve the infection and underwent an additional 2-month resolution. The prostate tissues were used for bacterial culture and measurement of hydroxyproline level as collagen content using high-pressure liquid chromatography.

Results: The uropathogenic E-coli was present in the prostates of all E-coli infected animals after 1-month post-inflammation and treatment of Baytril completely resolved the bacterial infection. We further found that hydroxyproline content in the prostates was significantly increased in the E-coli infected mice compared to the saline. Infected mice treated with Baytril with an additional 2-month resolution had a significant increase in hydroxyproline level in the prostates compared to the saline, but the level was significantly lowered to the E-coli infected prostates.

Conclusion: This study suggests that prostate fibrosis in response to bacterial-induced chronic inflammation is only partially reversed. Given that prostate fibrosis is suggested to strongly associate with the loss of prostate compliance and the development and progression of BPH/LUTS, an understanding on the mechanisms of the irreversible collagen deposition is required for improvement of therapeutic treatment.

252 Topical Application of TRPV1 Antagonist Effectively Inhibits PTD-Induced Sensitization in Murine Skin.

Transient receptor potential vanilloid type 1 (TRPV1) is expressed in the skin and plays a role in migration of dendritic cells to lymph nodes in allergic diseases. However, it has been unaddressed if TRPV1 blockade can suppress chemical-induced contact dermatitis. 1,4-toluenediamine (PTD) and 1,4-phenylenediamine (PPD) are widely used as permanent hair dye but repeated use of PTD can induce contact hypersensitivity by the induction of regulatory T cells in the draining lymph node and of inflammation in the exposed skin. In this study, the inhibitor effect TRPV1 antagonist, PAC-14028 on PTD-induced dermatitis and dendritic cell (DC) function was explored. This study was performed according to the method of nonradioactive local lymph node assay using flow cytometry. Groups of mice (N=4-5) were treated with 25 ul of the PTD alone, PAC-14028 alone, PTD and PAC-14028 or vehicle alone on the dorsal area of both ears daily for 3 consecutive days (Day 1, 2 and 3) and were sacrificed at Day 6. The local response was measured by ear swelling and by histological examinations (HE staining). We also used immunohistochemical staining western blot assay to evaluate the activation of specific inflammation makers and key mediators of signaling pathway in the mouse skin. The DCs migration markers in the draining lymph nodes were analysed by flow cytometry. TRPV1 antagonist suppressed PTD-induced edema in skin and proliferation of lymphocytes and inhibits the migration of CD11c+ DC and CD207+ Langerhans cells (LC) to the draining lymph nodes (LN).
Allergic airway disease (AAD) involves a complex interaction between various cell types within the lung, including inflammatory and epithelial cells. The inherent inflammation of AAD also leads to localized hypoxia. Communication between inflammatory cells, such as macrophages and T cells, and the epithelia in these hypoxic conditions is hypothesized to be critical to the progression of AAD. To characterize the role of epithelial-mediated hypoxia-induced signaling in AAD, epithelial and lung-specific conditional hypoxia inducible factor-1α (HIF1α), HIF2α, and HIF1α and HIF2α (HIF1/2α) knockout mouse models were created. Previous research in our lab has shown that the HIF1α-deficient (HIF1α/Δα) mice exhibit an exacerbated response to the ovalbumin (OVA) model of AAD. To determine the role of HIF2α in AAD and characterize possible compensation between the two HIFs following OVA challenge, recombination was induced early in postnatal development and HIF-deficient mice were then sensitized/challenged with OVA via a standard paradigm. In contrast to the HIF1α/Δα model, the HIF2α/Δα mice displayed no increase in eosinophil infiltration, T helper 2 cytokines, or airway resistance following OVA treatment. The HIF2α/Δα mice appeared phenotypically identical to OVA-treated littermate controls. Interestingly, when HIF1/2α/Δα mice were sensitized/challenged with OVA, these animals appeared similar to HIF2α/Δα and littermate controls, suggesting that HIF2α plays a role in the exacerbated inflammation observed in the HIF1α/Δα model. These data suggest that epithelial-derived HIF signaling plays an important role in establishing the immunity of the lung and that a proper balance between the two HIFs is required for a normal inflammatory response. The observed changes in the inflammatory response of the various models also suggest that early life exposures that alter the expression or function of HIF1α and/or HIF2α might have profound effects on the lung’s response to toxicant challenge upon reaching adulthood.

Environmental or occupational exposure to silica particles over an extended period of time is associated with the development of progressive inflammation and silicosis. Silicosis remains a prevalent health problem throughout the world. Currently, treatment choices for silicosis are limited and at present there is no cure for the disease. Therefore, it is essential to investigate potential therapeutic agents in silicosis treatment. Imipramine (IMP) is an approved tricyclic antidepressant, and a lysosomotropic agent. The aim of this study was to determine the protective effect of IMP on silica-induced inflammation and determine the mechanism by which IMP inhibits inflammation. C57BL/6 wild-type (WT) mice were used throughout the study. The protective effect of IMP was evaluated in vitro and 24 hours and 42 days following silica exposure in vivo. Silica was administered once a week for 4 weeks and IMP was delivered for 42 days via osmotic pumps implanted subcutaneously. Collagen levels were determined by hydroxyproline content. IMP treatment decreased collagen levels compared to the silica-exposed group. Lung histopathology improved on IMP treatment. WT mice were pretreated with IMP 25 mg/kg, and subsequently exposed to silica 1mg/kg during the in vivo acute study. IMP inhibited silica-exposed neutrophil infiltration, IL-1β and IL-6 in the lavage fluid. For in vitro experiments, WT alveolar macrophages (AM) were pretreated with 25 μM IMP and subsequently exposed to LPS (20 ng/ml) and silica (100 μg/ml). IMP was highly effective in blocking silica-induced inflammatory and toxic effects. The effect of IMP on acid sphingomyelinase (ASMase) was determined since A-SMase has been associated with acute lung injury. A-SMase was significantly inhibited by IMP treatment. The results demonstrate that IMP inhibits silica-induced inflammation and IMP may be exhibiting anti-inflammatory properties through its effect on lysosomes. The work was supported by NIH grants P20 RR017670 and R01 ES 15294.
and J. J. on potentiating amyloid pathology in AD. κβTLR-2 and microglia after PM2.5 exposure compared to filtered air. The results of transgenic animals, the hippocampus region showed enhanced staining for both β40 and Aβ activation, and IL-1β increased only in transgenic mice exposed to PM2.5. In transgenic animals, the hippocampus region showed enhanced staining for both TLR-2 and microglia after PM2.5 exposure compared to filtered air. The results of our pilot study suggest PM2.5 exposure may lead to the activation of innate immune response mediated by NF-kB pathway. The upregulation of β levels in the brain of PM2.5 exposed animals suggest that there could be a link between PM2.5 exposure and enhancement of amyloid production. The results of this pilot project justify the need for a more extensive study assessing the effect of particulate matter on potentiating amyloid pathology in AD.

Subchronic Exposure to Deltamethrin Causes Hippocampal Neuroinflammation and Deficits in Learning and Memory.

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Previously, we have reported that in vitro exposure of neuroblastoma cells to deltamethrin causes apoptosis through the ER stress pathway, leading to calpain and caspase-3 activation (Hossain and Richardson, 2011). Others have found that acute high-dose deltamethrin (12.5 mg/kg) exposure causes hippocampal apoptosis in adult rats (Wu and Liu, 2000). However, little is known about the effects of longer-term lower level exposure that does not result in acute poisoning. Here, we investigated the effects of deltamethrin at a dose of 3mg/kg every 3 days for 2 months on hippocampal ER stress, neuroinflammation, and learning and memory in adult mice. Deltamethrin treatment did not result in over toxicity. However, we observed increased spectrin cleavage in the hippocampus of deltamethrin-treated mice, indicating activation of calpain. Deltamethrin also significantly increased the hippocampal mRNA expression of glial fibrillary acidic protein (GFAP; 23%) and the pro-inflammatory cytokines tumor necrosis factor-alpha (121%) and Interleukin-1 alpha (96%), indicating an ongoing neuroinflammatory process. Finally, we found that subchronic deltamethrin exposure causes profound deficits in hippocampal-dependent learning and memory in the Morris water maze, which was accompanied by a decreased mRNA expression (30%) and protein level (39%) of nerve growth factor (NGF), respectively. Together, these data suggest activation of the ER stress pathway and neuroinflammation by repeated exposure to deltamethrin may contribute to the down-regulation of NGF in the hippocampus, which may result in subsequent impairment in learning and memory in mice. Supported by NIH ES015991 and ES005022.
Using a Sequence Homology-Based Predictive Strategy to Address Current Demands for Focused Toxicity Testing in Ecological Risk Assessment.


The lack of resources available for comprehensive toxicity testing, international interest in limiting the quantity of animals used in testing, and a mounting list of anthropogenic chemicals produced worldwide have led to the exploration of innovative means for identifying chemicals that are potentially hazardous to the environment and its inhabitants. Predictive toxicological approaches, which utilize publically available, a priori, knowledge of a chemical and its known molecular, cellular, or whole organism interactions, show promise for focusing current toxicity testing strategies. Using modern bioinformatic techniques, we have created a computational tool, which mines the extensive genomic and proteomic sequence repositories available through the National Center for Biotechnology Information and strategically compares homology metrics associated with primary and secondary protein sequences/structural domains across taxa. These comparisons are used to identify and rank species most likely to be susceptible to a chemical acting through a known molecular initiating event and can therefore aid in designing toxicity studies for species of concern. This presentation will identify the domains of applicability for this tool and describe examples related to predicting species sensitivities to pharmaceuticals and pesticides. An assessment of honey bee sensitivity to various pesticides will demonstrate the applicability of this tool for existing questions in risk assessment. The contents of this abstract neither constitute nor reflect official US EPA policy.

Alteration in Behaviour, Histoarchitecture of Liver, Lung, and Kidney of Male Wistar Rats Exposed to Open Refuse Dump.

P. U. Nwoha1, 2, 3, G. E. Waritimi2, A. D. Atomi2, F. Onyije2 and O. M. Ijjomene1.

Household wastes are disposed in the open in Nigeria, and most third world countries. These could have serious consequences on the health of humans and animals in the vicinity. Yet not much is known on the effects of such exposure on behaviour, and organs of the body. The present work exposed weaned male Wistar rats to open refuse dump continuously for five months. During this period the rats were housed in a building on the dump, kept in clean plastic cages, fed normal rat Chow, and provided clean drinking water. Control rats were housed in Niger Delta University Animal Holding. At the end of the period, behaviour of the rats was tested on Open, Elevated plus maze, for exploratory activity and for anxiety respectively. The animals were sacrificed, liver, lung, kidney dissected, examined macroscopically, and prepared for histological examination with H & E staining. Data obtained and analyzed with student t-test showed that rats exposed to refuse dump spent significantly less time in open arms of the elevated plus maze than their unexposed controls (p < 0.05). Organs showed massive infiltration with fat, while the histology showed destruction of the radial liver architecture, thinnning and collapse of alveolar wall of lung, and tubular wall of the kidney. Thus indicating that such exposure could induce anxiety, and massive destruction of organs of the body, leading to serious health consequences.

Key words: refuse exposure, anxiety, destruction, liver, lung, kidney

In Vitro Fish Metabolism Using Rainbow Trout Liver S9 Fractions to Evaluate the Bioaccumulation Potential of Fragrance Ingredients.


Bioaccumulation in aquatic species is a critical endpoint in the evaluation of novel chemicals as part of PBT assessment (persistent, bioaccumulative, and toxic) by the U.S. EPA. In vivo determination of the bioconcentration factor (BCF) requires the use of large numbers of animals. Predictive models are commonly used if no in vivo BCF data are available. These models generally acknowledge the possibility that biotransformation can reduce the extent of accumulation. Lacking measured data, however, modeled biotransformation rates are commonly set equal to zero. In vitro systems have been proposed as alternative methods that can be used to provide metabolic data needed to refine BCF computer model estimates.

The goal of our study was to determine the in vitro metabolic stability of common chemical classes of fragrance ingredients (esters, alcohols and ketones) using rainbow trout liver S9 fractions and to use in vitro clearance rate to model the bioaccumulation potential. Metabolic stability was determined by monitoring the disappearance of the parent molecule by GC MS and metabolite formation by GC-MS and LC-MS.

In vitro metabolic turnover was found with isosigmafilanolone (CAS 23787-90-8). The 5-carbon sesquiterpene (77218-42-1) was transformed rapidly. Alcohols like Ambermax (929625-08-1) and ketones like spirogalbanone (224031-70-3) were transformed moderately to rapidly. Metabolic routes, identified by selective use of cofactors and metabolite identification, involve ester cleavage, hydroxylations, reductions and conjugation with glucuronic acid and glutathione. When clearance rates measured in vitro were used as inputs to the BCF model a good correlation was observed between predicted BCFs and measured in vivo values. In vitro S9 metabolism data in combination with new refined BCF models are a valuable tool to assess bioaccumulation potential as part of a weight-of-evidence approach for chemical registrations.

The Role of Aquaporin 3 in the Uptake of Arsenite through the Intestine of the Atlantic Killifish (Fundulus heteroclitus).

D. Jung1, 2, M. A. Adamo2, R. M. Lehman2, R. Barnaby1, B. P. Jackson3, J. R. Shaw1, 2 and B. A. Stanton1, 2.

Aquaglyceroporins (AQPs) are proteins that mediate movement of water and small solutes across cellular membrane. Previously, we cloned kIAQP3a from the gill of the killifish (Fundulus heteroclitus), an environmental sentinel species. kIAQP3a, the only AQP expressed in gill, is the first AQP described that does not transport arsenite. This finding accounts for the low levels of cellular arsenite in gill of killifish exposed to environmental arsenite. Another homolog of AQP3 (kIAQP3b), which transports arsenic, was identified as the consensus from a transcriptome database. In this study, we sought to identify the AQP3s in the intestine, a major route of arsenite uptake. First, we examined AQP mRNA expression by qRT-PCR in the kilifish intestine. Among the AQP3s examined, only AQP3 was significantly expressed above background levels. Western blot studies with a polyclonal antibody that did not discriminate among kIAQP3 variants, revealed that kIAQP3 abundance was higher in killifish acclimated to FW compared to SW. Intriguingly, whereas only kIAQP3a was expressed in the intestine of FW killifish, both kIAQP3a and a new variant, kIAQP3c, were expressed in the intestine of SW fish. When kIAQP3c was transfected into HEK293T cells, cells took up arsenic as effectively as cells transfected with kIAQP3b. When we examined arsenic levels in the intestine of FW fish and SW fish exposed to 1000 μg/L arsenite for 72h, the amount of arsenic detected in the intestine of SW fish was higher than the amount detected in FW fish. Results indicate in the kilifish indicate that arsenite is most likely occurs via ingestion, and that killifish acclimated to SW take up more arsenite than FW acclimated fish, because kIAQP3c expression is up-regulated in SW fish and because SW fish drink more water than FW fish.

Characterization of DCOIT Bioaccumulation Mechanism via In Vitro Incubation with Rainbow Trout Liver S9.


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The goal of our study was to determine the in vitro metabolic stability of common chemical classes of fragrance ingredients (esters, alcohols and ketones) using rainbow trout liver S9 fractions and to use in vitro clearance rate to model the bioaccumulation potential. Metabolic stability was determined by monitoring the disappearance of the parent molecule by GC MS and metabolite formation by GC-MS and LC-MS.

In vitro metabolic turnover was found with isosigmafilanolone (CAS 23787-90-8). The 5-carbon sesquiterpene (77218-42-1) was transformed rapidly. Alcohols like Ambermax (929625-08-1) and ketones like spirogalbanone (224031-70-3) were transformed moderately to rapidly. Metabolic routes, identified by selective use of cofactors and metabolite identification, involve ester cleavage, hydroxylations, reductions and conjugation with glucuronic acid and glutathione. When clearance rates measured in vitro were used as inputs to the BCF model a good correlation was observed between predicted BCFs and measured in vivo values. In vitro S9 metabolism data in combination with new refined BCF models are a valuable tool to assess bioaccumulation potential as part of a weight-of-evidence approach for chemical registrations.
surrogate) in the presence or absence of NADPH. The potential metabolites including glutathione conjugates were identified and quantified via LC/MS-MS methodologies and radioactivity counting, and the protein binding activity of DCOIT was also quantified. Overall, more than nineteen metabolites (all chlorine-free) were identified with a majority of these metabolites conjugated to glutathione or cysteine. Incubations containing Fish S9 and DCOIT in the absence of glutathione resulted in much higher protein binding than those incubations containing glutathione. This demonstrated that glutathione present in the incubation medium acts as a useful surrogate of endogenous Fish S9 proteins to competitively react with DCOIT (or DCOIT active metabolites). The results from the present study provide evidence that the BCF value measured in vivo can be attributed to ring-opened metabolites binding to high molecular weight tissue fractions—likely the sulphydryl moieties in proteins. These data confirm that parent DCOIT does not bioaccumulate in fish, but the parent molecule is extensively metabolized to metabolites which were bond to high molecular weight components of the fish tissue (i.e., proteins).

270 Zinc Content Determines the Toxicity of Tire Leachate in Girardia tigrina.


The practice of recycling old tires into various outdoor structures such as play-ground surfaces and landfill liners poses the risk of tire components moving into local waterways and possibly affecting aquatic organisms. In this study we tested the hypothesis that the zinc content of tire leachate is a significant factor in its toxicity to Girardia tigrina (Girard, 1850), a freshwater planarian common to North American waterways. Planarians were cultured in tire leachate containing either 49.5 mg/L of zinc (BALT) or 0.13 mg/L (FRESH) of zinc or a control of extraction medium (EM) over a time period of 24 hours. All planarians in the BALT group died within 24 hours while no planarians died in either the FRESH or EM group. To verify that zinc was the causative agent in the observed toxicity, planarians were maintained in a solution containing an equivalent amount of zinc (from zinc sulfate) for 24 hours. The survival rate of planarians in this group was not significantly different from the survival rate observed in the BALT group. These data strongly indicate zinc as the toxic agent. In addition to the lethality demonstrated by both high-zinc solutions, planarians displayed signs of distress indicated by increased activity, writhing, and loss of motor coordination prior to death when compared to controls, suggesting that neurotoxicity may be the mechanism of action. Future studies will examine the dose-response relationship of zinc toxicity in G. tigrina as well as evidence of neurotoxicity. (Support: NSF Award 0928444.)

271 Harmonizing Use of the 3Rs in Fish Toxicity Testing.

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Pursuit of methods that refine, reduce or replace animals is often discussed in the context of human health hazard and risk assessment; however, several approaches that are in common use for human health are more frequently being applied in ecological hazard and risk assessment as well. Organized testing frameworks such as tiered frameworks or integrated strategies can be used to prioritize information needs and focus testing on tests that would be the most informative for a given regulatory need. Specific application of integrated strategies has been used to minimize fish used for acute toxicity testing, for example the limit and threshold approaches. There are principles that can be applied to specific test protocols that minimize animal use, for example the use of historical controls where appropriate or using statistical analyses to define the minimum individuals needed to obtain statistically significant results. Computer modeling can assist in extrapolating information from one species to another or in predicting acute toxicity, bioaccumulation and other biological activity in fish. While limited embryo, ex-vivo, and in vitro approaches are currently used in fish toxicity testing, opportunities exist for expanding the repertoire, particularly in the area of ‘omics technologies. This presentation will describe these approaches as currently applied and present recommendations for improving the application and harmonization of 3Rs approaches to fish toxicity testing, as presented in Chapter 5 of the Fish Toxicity Testing Framework Guidance Document prepared by the Organization of Economic Cooperation and Development (August 2012).

272 Concentrations of Metals Associated with Crude Oil from the BP Macondo Well in Sediments and Fish from the Northeastern Gulf of Mexico.

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To investigate if metals associated with crude oil from the BP Macondo Well were entering the marine food chain, sediment and scad mackerel (TRACHURUS LATHAMI) samples were analyzed by ICP for five of these metals: chromium, nickel, lead, thallium and vanadium. Samples were collected from the carbonate shelf along the west coast of Florida in fall, 2010. A subset of these samples was collected from the western Florida Panhandle. This area was south of the Florida beaches where tar balls washed ashore during the summer of 2010. Nickel levels in samples from this subset ranged from 2.85 to 11.18 micrograms/g in dry sediments, and 0.00 to 0.86 micrograms/g in dry fish tissues. Chromium levels in samples from this area ranged from 7.08 to 12.43 micrograms/g in sediments, and from 0.00 to 0.76 micrograms/g in scad. Vanadium levels ranged from 3.32 to 10.07 micrograms/g in sediments. Vanadium was the only one of these metals not detected in any of the fish. Lead levels ranged from 1.13 to 5.89 micrograms/g in sediments, and 0.00 to 0.36 micrograms/g in scad. Thallium appears to be biomineralized in scad, as concentrations in fish ranged from 0.72 to 1.42 micrograms/g, while sediment levels ranged from 0.00 to 0.30 micrograms/g.

273 The Effect of Tributyltin (TBT) on Zebrafish Sexual Differentiation.

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Tributyltin (TBT), an antifouling agent, has been implicated in the masculinization of fish species worldwide, however the molecular mechanism is not fully understood. Our lab has previously examined the actions of TBT as an endocrine disruptor in zebrafish (Danio rerio) and determined, in vitro, that TBT inhibits zER-specific activity in a dose dependent manner and may potentially act through the RXR portion of the PPAR-RXR (peroxiome proliferator-activated receptor gamma-retinoid X receptor alpha) heterodimer. Additionally, zebrafish were exposed to increasing concentrations of TBT and sexual differentiation genes were analyzed via qPCR. Results from qPCR focused our experimental efforts on the candidate gene, SRY-box containing gene 9α (Sox9α). Sox9α is a key regulator in mammalian testis differentiation, where it is shuttled to the nucleus upon differentiation; this appears to be a conserved mechanism across fish, marsupials and placental mammals. Developmental toxicity embryonic embryos were exposed to 1pM and 2.5pM TBT from 10 days post hatch (dph) to 90 dph and sampling was done at 25, 35, 40, 45, 60 and 90 dph. Fish were treated three times per week with either TBT, estrogen or vehicle. Following treatments, immunohistochemical (IHC) analysis was performed to assess Sox9α nuclear or cytoplasmic localization.

274 Heavy Metal Response in Daphnia magna: An Ecologically- Relevant Nonmodel Organism.

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Environmental health issues have become a major focus of ecotoxicology research over the last decade. Due to rapid industrialization and urbanization, ecosystems are currently, and into the foreseeable future, under the threat of potential damage. In order to mitigate these risks, environmental scientists build predictive models to gauge the impact of pollutants, perform risk assessment studies to measure water quality, and protect organisms from potential damage due to the effects of pollutants. One major pollutant is heavy metal. Since aquatic systems are the major sinks of industrial effluents they are often highly impacted by heavy metals than area terrestrial ecosystems. Numerous studies on individual organisms and populations have demonstrated the effects of acute exposure to metal pollutants, but few studies are available that show the consequences of long-term low dose metal exposure to aquatic organisms. In regular risk assessment practice, acute and high dose/concentration exposure is a common approach to predict which metal/chemical has potentially harmful effects on a particular organism. This approach is informative, but not predictive of the long-term low dose/concentration scenario that is more likely happening in our
day-to-day life. Further, aquatic toxicity tests have historically been largely limited to model organisms. However, not all organisms show the same physiological response to a particular metal/chemical making these assessments species specific. Our approach is to use quantitative genetics/genomics tools applied in a ecological relevant non-model organism to determine the genetic basis and mechanisms of response to a common, toxic metal pollutant. We use Daphnia magna, a widespread freshwater invertebrate, as a model organism to understand and interpret the genetic mechanisms of response to cadmium.

**275 Characterization of the Hepatic Metabolome of Migrating Sockeye Salmon in British Columbia, Canada.**

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The health of British Columbia wild sockeye salmon (Oncorhynchus nerka) is of increasing concern due to recent extreme variation in the number of fish returning to spawn annually. Causes of this variability are unclear but may be related to contaminant or viral exposures, climate change, or food shortages. Two key migratory routes for Pacific sockeye are the Fraser and Skeena River watersheds. These watersheds represent highly contrasting environments for spawning salmon; the former passes through populous and industrialized regions of Vancouver, while the latter flows through fairly remote regions with little industrial input. A recent comparison of hepatic mRNA profiles between Fraser and Skeena sockeye revealed stark differences in estrogen-associated signaling in Skeena fish, despite these fish spawn in a relatively pristine environment. In contrast, the status of hepatic gene transcripts for Fraser River sockeye showed normal reproduction-related changes in estrogen-associated signaling. To expand upon available toxicological endpoints and further define potential sources of exposure resulting in the observed hepatic mRNA profiles in Skeena River salmon, we compared aspects of the hepatic metabolome of fish from both populations. A total of 186 metabolites from 6 different metabolite classes were measured using a recently developed assay. Molecular targets included acylcarnitines (n=40), amino acids (n=21), glycerophospholipids (n=90), hexoses, sphingolipids (n=15), and biogenic amines (n=19). Major metabolites quantified in Sockeye salmon liver included glycine, carnitine, phosphatidylcholine acyl-alkyl C38:6, and sphingomyelins C24:1. The combined transcriptomic and metabolomic data provides a comprehensive molecular profile of fish health, and sheds light on the biochemical changes arising from alteration of estrogen-associated signaling.

**276 Elevated Metals and Organic Concentrations Linked to Biomarkers’ Alterations in Organisms Exposed to Mining Effluent.**

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The Blesbokspruit wetland, South Africa, continuously receives diffuse and point source releases of mining effluent till date. Results are presented from a first time study conducted in the system on metals and organic bio-accumulation in sediments, residents and transplanted organisms. A suite of biomarkers assay was also carried out in bio-indicators. During the 2008 low and high flow periods, resident catfish, Clarias gariepinus and tilapia, Tilapia sparrmanii species were collected from 5 sites in a field survey. During the 2009 high flow, a transection study (active bio-monitoring - ABM) was conducted using laboratory reared Tilapia spp. at four of the five sites in cages for four weeks. Biomarkers of exposure: cytochrome P-450 (CYT-P450) and acetylcholine esterase (ACHE) and biomarkers of effect: catalase (CAT) and superoxide dismutase (SOD) responses were determined in all samples. Metal and organic concentrations measured varied among sites and were elevated (p<0.05) during the field survey high flow; this was the case in most cases. This was always high for both periods in Site 1 which is upstream and close to mine dumping. Biomarkers were found altered both in resident and transplanted organisms at all sites. For example, triggered CAT activity and inhibition of SOD both indicative of oxidative stress was observed. CYT-P450 and ACHE activities inhibition, which indicates organometallic and pesticides exposure was also observed. Biomarker responses were similar for both resident and transplanted fishes for the high flow periods (p<0.05). Site 5 (reference site) generally showed the least altered responses during the field survey low flow but not during the high flow and ABM. Biomarkers were able to successfully demonstrate biological effects from toxicants in the system. It was possible to link these effects to observed elevated metal and organic concentrations found in the bio-indicators which can be applied within the proposed integrated management plan for the Blesbokspruit as well as other catchments.

**277 Effects of Two Progestins, Norethindrone and Levonorgestrel, on Reproduction in a Marine Fish, Taurogadus adspersus.**

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Endocrine-active pharmaceuticals that enter the aquatic environment through sewage effluent may have unintended impacts on reproduction in fish, which in turn may affect the sustainability of exposed populations. Laboratory experiments were conducted with the marine fish cunner (Taurogadus adspersus) to evaluate whether norethindrone (NOR) and levonorgestrel (LNG) affected reproduction in spawning adults. Both progestins are used in human contraceptive formulations and have been detected in low (ng/L) concentrations in aquatic environments. Synthetic progestins in aquatic environments are of special concern because some fish use natural progesterones as phenotypes to coordinate reproduction, and evidence suggests progestins may be selectively taken up through the gills in some species. Reproductive endpoints of egg production, viability and fertility were assessed daily in spawning cunner treated with NOR or LNG (nominal concentrations of 0.0075 or 0.75 mg/kg) by oral gavage on days 0, 4, 8, 12 and 16 of the experiment. All fish were sacrificed on day 17 and gonadosomatic index (GSI) was determined. In NOR-treated fish, egg production per gram female was significantly reduced relative to controls at both concentrations, while egg fertility and viability was notably decreased, although not significantly, only in the 0.75 mg/kg treatment. GSI was significantly reduced in both males and females from the 0.75 mg/kg treatment. Female mortality in this treatment group was more than twice that in controls, indicating an increase in male aggression. In LNG-treated cunner, no significant effect was seen on egg production, fertility, viability, or GSI compared to control fish. Results indicate some progestins can impact fish reproduction, even in short-term exposures. Research is planned to determine if these fish selectively take up progestins from the aquatic environment. This abstract does not reflect US EPA policy.

**278 Effects of Fungicides on Honey Bee Development and Behavior.**

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Pesticides may contribute to the health challenges facing honey bees. Bees experience chronic exposures through contaminated beeswax and stored pollen and honey. There is a need to study possible sublethal or delayed effects from such exposures, which may ultimately result in the collapse of the colony. Fungicides are thought to have little effect on insects, and are routinely sprayed while bees are pollinating crops. Multiple fungicides are transported with pollen into the colony, and some are known to persist in beeswax. Beekeepers suspect that fungicides may have an effect on honey bee development, and some laboratory tests have demonstrated adverse effects of fungicides on bee larvae. We have developed laboratory methods to chronically expose young adult bees to pesticide-contaminated beeswax, in concentrations similar to those found in hives. By using Noldus EthoVision to track the behavior of bees on video, we have found that the major contaminants of beeswax, including the fungicide chlorothalonil, delay behavioral development as measured by the initiation of circadian activity rhythms. Young adult bees consume pollen and secrete proteinaceous brood food and royal jelly to feed developing larva and the queen. In semi-field experiments, we fed pollen spiked with fungicides to colonies of bees, similar to field concentrations. By evaluating colonies weekly, we found that chlorothalonil and iprodione affect larval development, and ziram affects queen health several weeks after initial exposure. These results suggest that chronic contact exposure through wax and ingestion of fungicides through pollen may target the development and social function of young worker bees, and may have detrimental effects on the colony.

**279 The Mediterranean Gecko, Hemidactylus turcicus (Gekkonidae : Squamata)—An Alternative Model for the Study of Redox Potential.**

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Glutathione and similar sulphydryl groups play an important role in redox cycling in mammals. In this study we have investigated the role of sulphydrls in the Mediterranean Gecko, Hemidactylus turcicus (Gekkonidae : Squamata). Concentrations of hepatic sulphydrls were determined in the field controls and the geckos maintained at various temperatures. Concentrations of sulphydrls in livers
of field controls were 46.3 ± 5 mmol/L and were significantly elevated (141 and 151 % of controls) in geckos maintained at 15 and 10°C, respectively. Female geckos had significantly higher (139 %) concentrations of sulfhydryls than did the males. This study indicates that this oxidative biochemical pathway is operative in geckos. Geckos may provide a very inexpensive alternative animal model for redox studies.

280 Recent Emergence of Perfluorohexanoate in Tap, River, and Sea Water in Japan.
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Environmental waters such as river water (RW) and sea water (SW) are expected to be the major exposure sources of Perfluorooalkyl acids (PFAA) to humans via tap water (TW) and food fish. At the 2012 Annual Meeting of SOT, we reported the trend in PFAA contaminations in Japanese RW and TW from 2003 to 2010, based on the results of measurement of Perfluorocarboxylates (from C5 to C12 carbon backbone) and perfluorosulfonates (CF2, C6F2, C8F2 and C10F2). The major PFAA (C6, C8, C9 and Cs8) concentrations in both RW and TW were always highest in Kinki area. In 2010 extremely high RW C6 concentrations (46 and 24 μg/L) were detected in the lower reaches at the foot of a fluorochemical plant, where extremely high concentrations of C8 (67 and 24μg/L) had been detected in 2003. From 2003 to 2010, Kinki showed drastic reduction of RW C8 concentration to one tenth. C6 concentration in TW in 2007 showed as low as 2.3 % of RW in 2010 compared to 42.3% for that of C8. The conclusion was that the release of C6 to the environment had begun recently from the source in Kinki. In the present study, we measured PFAA (from C4 to C16; and C4S, C6S, C8S and Cs10) in RW (12 locations) and TW (6 locations) in the Kinki area and coastal SW around Japan (31 locations) collected in 2011 using LC-MS/MS. The highest RW Perfluorocarboxylates (from C4 to C10) were detected in the lower reaches at the foot of the fluorochemical plant (for C6: 49 and 43μg/L). The highest RW C6 concentration in the 2011 samples was 2.85 μg/L, which was greater than the highest C6 concentration (1.51 ng/L) in the 2007 samples. The highest SW C6 concentration in Kinki was far greater than the samples from the other areas and was 129 ng/L. From these results it was concluded that C6 release from the source to the river of recent onset is rapidly contaminating surrounding SW and gradually contaminating TW in the nearby areas.

281 Distributions of Metals (Cadmium, Lead, Iron, Manganese, Zinc, and Copper) in Water, Aquatic Plant, and Fish.
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Concentrations of cadmium (Cd), lead (Pb), ferrum (Fe), manganese (Mn), zinc (Zn) and copper (Cu) were measured in water, Ceratophyllum demersum (C. demersum) aquatic plant, Clarias lazera fish (C. lazera) collected from nine sampling stations along El Ebrahimia canal and two districts located at the east bank of the Nile in the province of Beni Suef, Egypt during 2009-2010. Atomic Absorption analysis revealed that the studied metals were higher than the limit of detection (LOD) in all the examined samples. In water, Pb had the highest concentration among the metals detected in seven of the nine tested locations (0.3 - 0.9 ppm). The concentrations of Pb, Fe, and Mn were above the maximum Egyptian permitted limits in all tested sites, while Zn and Cu concentrations were below the permitted limits (4 mg/L and 2 mg/L) in the nine districts. Comparisons were made of the metal concentrations in water and aquatic plants with those in the catfish tissues obtained from water. The metal concentrations found in the C. demersum aquatic plant samples taken in the nine studied districts were distributed in this order; Mn > Zn > Cu > Pb > Fe > Cd. and were higher than the water In fish, metals accumulated in the various examined tissues at several levels, but the metal concentrations in muscles (edible part) were below the metal levels in the other organs (nonedible) in the fish samples. The concentrations of Cd, Pb and Fe in fish tissues were above the international standard, while the concentrations of Mn, Zn and Cu were below this standard. The high concentrations of these metals in water, aquatic plants and fish in El Ebrahimia canal may be the result of both anthropogenic activities producing industrial, agricultural and domestic waste and accidental pollution incidents.

282 Cytochrome P450 Monoxygenases Expression in Human Epithelial Lung Cell Lines.
M. Niehof and T. Hansen. Fraunhofer ITEM, Hannover, Germany. Sponsor: C. Dassenbrock

The pulmonary epithelium is the first barrier for airborne xenobiotics and inhaled drugs. Cytochrome P450 monoxygenases (CYP) participate in metabolic inactivation of xenobiotics. Beyond that some compounds require enzymatic activation to exert their toxic effects or their desirable functions. The bronchiolar epithelial cell line Calu-3 and the type II-like pulmonary epithelial cell line A549 serve as cell culture models for the human respiratory epithelium. So far, there is little information regarding their metabolic properties especially for Calu-3 cells. The goal of this study was to further characterize both cell lines regarding their basal and inducible CYP isoform expression.

CYP expression was determined using real-time reverse transcription quantitative polymerase reaction (RT-qPCR). Basal expression of CYP1B1, CYP1A1, CYP2D6, CYP2B6, CYP3A5, and CYP2J2, and slight amounts of further CYPs were detected in both cell lines, which is consistent with expression in the human lung.

Furthermore, potential CYP inducers were analyzed. Omeprazole acts on anly hydrocarbon receptor (AhR) activation and induced CYP1A1 and CYP1B1 in both cell lines. Induction acts on p53, p63 and PXR and the glucocorticoids act predominantly on constitutive androstane receptor (CAR), however, both receptors are not expressed in the lung. Accordingly, both agents did not induce any CYP in these cells. Besides PXR, dexamethasone acts on glucocorticoid receptors and we found induction of members of the CYP3A family, mainly CYP3A7 in Calu-3 cells, and CYP3A5 and CYP3A7 in A549 cells. CITCO is known to act as a CAR agonist and is normally used to induce CYP2B6/7. However, it is a potent inducer of CYP1B1 in Calu-3 cells, and of CYP1A1 and CYP1B1 in A549 cells. Thus, Calu-3 cells and A549 cells express a broad range of CYPs with preserved inducibility and are valuable models of the airway epithelial barrier for metabolic in vitro experiments.

283 Role of Renal Proximal Tubule P450 Enzymes in Chloroform-Induced Nephrotoxicity.
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The kidney is a primary target for numerous toxicants. Cytochrome P450 enzymes (P450s), responsible for the metabolic activation of various chemical compounds, are predominantly expressed in proximal tubules in the kidney. However, the specific role of proximal tubule P450s in chemical-induced nephrotoxicity is unclear. The aim of this study was to test the hypothesis that renal proximal tubule P450s are critical for the metabolic activation and nephrotoxicity of chloroform. To test this hypothesis, we have developed two new mouse models, one having proximal tubule-specific deletion of the cytochrome P450 reductase (Cpr) gene (the enzyme required for all microsomal P450 activities), named kidney-Cpr-null, and the other with proximal tubule-specific rescue of CPR activity in a model with global suppression of CPR activity in all extra-renal tissues, named extra-renal Cpr-null. The kidney-Cpr-null, extra-renal-Cpr-low, Cpr-low, and wild-type (WT) control mice were treated with a single oral dose of chloroform at 200 mg/kg. Blood, liver and kidney samples were obtained at 24 h after the treatment. Kidney toxicity was assessed by measuring serum levels of BUN and creatinine, and by pathological examination. The blood and tissue levels of chloroform were also determined. Chloroform-induced proximal tubular lesions and increases in BUN and creatinine levels were observed in all four genotypes, but the severity of toxicity was less in kidney-Cpr-null and Cpr-low mice, compared to WT and extra-renal-Cpr-null mice, respectively. There was no significant difference in chloroform levels in the blood, liver, or kidney, between kidney-Cpr-null and WT mice, or between extra-renal-Cpr-low and Cpr-low mice. These findings indicate that local P450 dependent metabolic activation plays an important role in renal toxicity induced by chloroform. Our results also demonstrate the utility of these novel mouse models for studies on the renal toxicity of other chemicals.

284 Green Tea Epigallocatechin Gallate Inhibits Drug Metabolizing Enzymes by Covalent-Binding to the Proteins and Formation of Protein Aggregates.

Green tea supplements have been reported to cause hepatotoxicity but the mechanisms of the toxic metabolite, epigallocatechin gallate (EGCG), are unknown. In this study, the rat liver microsomes were treated with 1 - 100 μM EGCG for 30
min, and the EGCG-binding proteins were affinity purified and probed with anti-

bodies against glyceraldehyde-3-phosphate dehydrogenase (GAPDH), actin, cy-
tochrome P450 (CYP) 1A1, CYP1A2, CYP2B1/2, CYP2E1, CYP3A, catechol-O-
methyltransferase (COMT) and microsomal glutathione transferase 1 (MGST1).
All but actin and soluble COMT were positively detected at ≥1 μM EGCG, indi-
cating EGCGs selectively bound to a subset of proteins including membrane-bound COMT. The binding correlated well with inhibition of CYP activities, except for CYP2E1 whose activity was unaffected despite evident binding. When microsomes were probed on Western Blots, all the actin and CYP2E1 antibodies showed a significant reduction in binding at ≥1 μM EGCG, suggesting that a fraction of the indicated proteins formed aggregates that were not recognizable by antibodies against the intact proteins. Protein aggregate formation was also observed in Coomassie Blue-stained SDS-PAGE gels. EGCG effects were partially abolished in the presence of 1 mM glutathione. We conclude that EGCG inhibits drug metab-
olizing enzymes by covalent-binding to the target proteins and formation of protein aggregates.

285 Metabolism of Rutacearpine in Freshly Isolated Hepatocytes from Rats and Mice.

T. Jeong, D. Lee, D. Oh, J. Kim, Y. Jahng and M. Kang, Pharmacy, Yeungnam University, Gyeongsan, Republic of Korea.

Rutacearpine is an alkaloid originally isolated from Evodia rutacearpa that has been used for the treatment of gastrointestinal disorders in Asia. In the present study, Phase I and Phase II metabolisms of rutacearpine were investigated in freshly iso-
lated hepatocytes from rats and mice. The results indicated that the metabolism of rutacearpine in rats and mice was different. When rutacearpine was incubated with fresh hepatocytes isolated from either rats or mice for 2 hr, 5 major Phase I metabolites were observed in both hepatocytes with different extents. Likewise, the pro-
duction of sulfate conjugates also showed difference. In rat hepatocytes, 4 sulfate conjugates were observed, whereas only two conjugates were observed in murine hepatocytes. The results indicated that the species selection would be concerned in the process of preclinical investigation Supported by a grant from National Research Foundation of Korea (2010-002666220).

286 Effects of Intestinal Microflora on Oral Pharmacokinetics of Baicalin in Normal and Antibiotic-Treated Mice.

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Baicalin (baicalein-7-glucuronide) is an ingredient of Scutellaria baicalensis Georgii that has been used as one of the most popular herbs in Korea for treatment of inflamma-
tion, cardiovascular diseases, hypertension, and microbial infections. In the present study, effects of intestinal microflora on oral pharmacokinetics of baicalin were investigated in normal SPF and antibiotic-treated mice. To control the number of intestinal bacteria, mice were pre-treated orally with erythromycin, oxytetracy-
cyclin and ceftadroxil for 3 consecutive days, followed by an oral administration with 100 mg/kg baicalin. Then baicalin and its possible metabolites in serum were determined using liquid chromatography/electrospray ionization mass spec-
trometry. By the pre-treatment with antibiotics, the number of intestinal microflora was significantly reduced. In addition, serum concentrations of baicalin and its metabolite, baicalin-6-glucuronide, were remarkably changed by treatment with antibiotics when compared with control mice. These results indicated that the in-
testinal microflora might have a critical role in modulating oral pharmacokinetics of baicalin. Supported by the grant from KFDA (09172KFDA996) and from National Research Foundation of Korea (2010-002666220).

287 Role of Intestinal Microflora in Oral Pharmacokinetics of Baicalin in a Germ-Free Mouse Model.

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Baicalin and its aglycone baicalein are bioactive flavonoids originally isolated from the root of Scutellaria baicalensis Georgii, a medicinal plant that has been used for the treatment of inflammation, hypertension, cardiovascular and allergic diseases. In the present study, role of intestinal microflora in baicalin metabolism was inves-
tigated following a single oral administration with 100 mg/kg baicalin in germ-free and control mice. Baicalin and its metabolites were determined by HPLC coupled with a tandem mass spectrometry. Serum concentrations of baicalin and its metabolite in germ-free animals were significantly lower than those in control mice having normal intestinal microflora. Likewise, transient hepatotoxicity induced by baicalein in germ-free mice was significantly different from that in control mice. These results indicated that the intestinal microflora might play a key role in the metabolism of baicalin orally ingested. Supported by the grant from KFDA (09172KFDA996) and from National Research Foundation of Korea (2010-002666220).

288 Importance of Chirality Considerations in Risk Assessments: Enantiomer-Specific Pharmaceutical Metabolism with Rainbow Trout (Oncorhynchus mykiss) S9.

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Enantiomers are capable of having significantly different biological effects, selectiv-
ity for receptors/transporters/enzymes, potency, and biodegradation rates. Differences in environmental fate, bioavailability and toxicity have also been re-
ported. Despite this knowledge, enantiomers are often treated as a single chemical entity in environmental monitoring and risk assessment. Enantiomer-specific dif-
fferences in pharmaceuticals are especially well documented within mammalian lit-
erature. Finding a way to leverage existing pharmaceutical safety data and pharma-
cology information through biological “read-across” may aid our ability to perform more accurate environmental assessments. In this study, we examined the compar-
tive metabolism of R, S and the racemate of three pharmaceuticals (propranolol, ibuprofen and fluoxetine) in rainbow trout liver S9 using a substrate depletion ap-
proach. An isotope dilution liquid chromatography tandem mass spectrometry (LC-MS/MS) method was employed for quantitation of parent chemical concen-
trations. Differential substrate depletion rates were observed for both R and S enan-
tiomers. The fastest clearance rates were observed with rac-propranolol, followed by S and R-propranolol, respectively. Ibuprofen appeared to undergo limited me-
tabolism; however, the resulting depletion curves did not differ statistically from those obtained for denatured controls. No substrate depletion was observed for rac, R or S fluoxetine. Mammalian clearance rates will be compared and risk assessment implications discussed.

289 The Role of Interstrain Differences in Trichloroethylene Metabolism in Kidney Effects in Mice.

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Trichloroethylene (TCE) is a well-known environmental and occupational toxicant contaminating air, water, and soil. The U.S. EPA recently issued the final IRIS as-
seessment of TCE and classified TCE as carcinogenic to humans. Still, several issues critical for assessing human health risks from TCE remain unresolved, such as (1) the amount of glutathione (GSH)-conjugated metabolites formed in various tis-
tures, and possible inter-individual and inter-species differences; and (2) the mode of action involved in kidney toxicity/carcinogenesis. The aim of this study was to use a panel of inbred mouse strains to investigate the relationship between inter-
strain differences in TCE metabolism and kidney toxicity. TCE (600 mg/kg/day, in 5% Alkamuls EL-620 in saline) was administered by gavage to male mice (6-8 weeks old) from 7 inbred strains (129S1/SvImJ, A/J, BTBR T+) for 5 days. Liver, kidney, and serum were collected at 2 and 8 hrs after the last dose. Quantification of S-(1,2-
dichlorovinyl)-L-cysteine (DCVC), S-(1,2-dichlorovinyl)-L-cysteine (DCVC), and trichloroethanol (TOCH) was performed. In addition, blood urea nitrogen, kidney-to-body weight ratio, proximal tubular cell proliferation, expression of per-
oxides proliferator marker genes (Ppara, Acox1, and Cyp4a10), kidney injury molecule-1 in kidney were evaluated. Inter-strain variability in the levels of DCVC, TCE, and TOCH in liver, serum, and kidney were observed. Overall, the level of TOCH was 10,000-fold greater than that of DCVC and DCVC in both liver and kidney. In conclusion, the inter-strain differences in TCE metabolism provide a mechanistic basis for examining the inter-strain differences in TCE organ-specific toxicity. This work was supported by the Superfund Basic Research Program grant P42 ES009548.
Chloroform (CF) is a tributylmethane present in drinking water as a byproduct of disinfection, and the kidney is one of the targets of toxicity in experimental animals. Since CF induces toxic effects via production of reactive metabolites, proper characterization of metabolism is essential for risk assessment. A revised physiologically-based pharmacokinetic (PBPK) model, in conjunction with benchmark dose (BMD) modeling, was used to interpret rat and mouse renal toxicity markers in three oral and inhalation studies. Large species differences in potency as a function of external dose became minimal when expressed in terms of a renal dose metric (daily mg CF metabolized per L cortex), indicating that species differences in susceptibility may be primarily mediated by toxicokinetics, rather than differences in toxicodynamics. The external BMD result for nuclear enlargement was 6 ppm for the mouse, but 40 ppm for the rat—a 7-fold difference. When the measure of dose was changed from inhaled concentration to the renal dose metric, the BMD result becomes 62 mg/L in the mouse, and 47 mg/L in the rat—a difference of 30%. Since results derived from drinking water data were also consistent with inhalation data, the work presented here increases confidence in the PBPK model, and the use of site-specific chloroform metabolism as the internal dose-metric. The views expressed in this publication are those of the authors and do not represent the views and policies of their respective Agencies.

**290** Interpretation of Multiroute Data for Chloroform-Induced Renal Toxicity in Rats and Mice Using an Updated Physiologically-Based Pharmacokinetic Model.

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N-Acetyltransferase 1 (NAT1) has similar toxicological profiles yet information on the metabolism of petrogenic PAHs is lacking. We report the metabolic fate of 1-methyl-phenanthrene and 9-ethyl-phenanthrene as representative alkylated petrogenic PAHs in human hepatoma (HepG2) cells. The structures of the metabolites were identified by HPLC-UV-fluorescence detection and LC-MS/MS. Both 1-methyl-phenanthrene and 9-ethyl-phenanthrene showed the formation of O-sulfated mono-phenols, O-sulfated bis-phenols, O-sulfated dihydrodiols, and O-sulfated catechols. The identification of these sulfate conjugates supports metabolic activation of 1-methyl-phenanthrene and 9-ethyl-phenanthrene by P450 and AKR isozymes followed by metabolic detoxification by SULT isozymes. (Supported by U19ES020676-01 to TMP).

**293** N-Acetyltransferase 1 (NAT1) Expression and Activity during Keratinocyte Differentiation and Cell Cycle Progression.

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N-Acetyltransferase 1 (NAT1) dependent N-acetylation is an important detoxification pathway for arylamines including certain dyes. Proliferating keratinocytes have high N-acetylation capacities, but the influence of differentiation on NAT1 is not clear.

Keratinocyte differentiation is associated with an arrest of cells in the G0/G1 phase of the cell cycle. In order to analyze NAT1 regulation in the different cell cycle phases we synchronized HaCaT keratinocytes by serum starvation (83±4% G0/G1) and re-addition for 20hrs (66±1.7% S-phase) as well as by double thymidine block (68±2% G0/G1, 72±6% S-Phase) and analyzed NAT1 activity, protein and mRNA expression. In line with the high N-acetylation capacity of the skin, NAT1 activity was compared to S-phase, elevated (about 40±2%) in the G0/G1 phase, which is the predominant state of keratinocytes in the epidermis. NAT1 protein levels were also higher in G0/G1 phase, while NAT1 promoter P1 dependent steady state mRNA levels were not enhanced.

In the next step, we differentiated the keratinocyte cell line HaCaT and primary keratinocytes in vitro and analyzed NAT1 mRNA expression and NAT1 activity. With increasing differentiation we found no NAT1 variation after in vitro differentiation and detected NAT1 protein staining throughout the entire epidermis using human skin slices.

These results indicate that in vitro differentiated keratinocytes do not lose N-acetylation capacity, although cell proliferation is terminated, possibly due to high NAT1 activities in G0/G1 phase arrested cells. However, variations of the keratinocyte cell cycle phase distribution may influence NAT1 activity and thereby detoxification capacities.

**294** Characterization of Peroxidase Activity in SkinEthic® Reconstructed Skin Models Compared with Ex Vivo Human Skin Samples.


Skin metabolism is becoming a major consideration in the development of new cosmetic ingredients, skin being the first organ exposed to them. Consequently, the use of ex vivo samples of normal human skin (NHS) or reconstructed human skin models (skin models) as alternative tools to animal testing requires to characterize and compare their abilities to metabolize xenobiotics. In this work, we determined if they possessed a functional peroxidase activity. Previous studies showed that NHS and skin models from SkinEthic® Laboratories such as Episkin® and the full thickness model of Episkin® expressed the mRNAs of several peroxidase isoforms (mainly in GPx and COX families). The catalytic activity of these enzymes was measured from dose-response studies using cumene hydroperoxide as substrate. Apparent Vmax, Km and ratio Vmax/Km (assessing metabolic rates) were comparable to the CYP3A4 micosomal control incubations. These results indicate that this system has the potential to improve ones ability to detoxify chemicals. Further studies are needed to determine the capability of the TNDs to metabolize chemicals in vivo.

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**292** Sulfonation of 1-Methyl-Phenanthrene and 9-Ethyl-Phenanthrene in Human Hepatoma (HepG2) Cells.

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Exposure to petrogenic polycyclic aromatic hydrocarbons (PAHs) in the food-chain is the major human health hazard associated with the Deepwater Horizon gulf-oil spill. Risk assessment is based on the assumption that petrogenic and pyrogenic PAHs have similar toxicological profiles yet information on the metabolism of petrogenic PAH is lacking. We report the metabolic fate of 1-methyl-phenanthrene and 9-ethyl-phenanthrene as representative alkylated petrogenic PAHs in human hepatoma (HepG2) cells.
Covalent protein adduction, which can underlie drug toxicity and/or reflect exposure, is largely unstudied in the case of illicit drugs of abuse. This research investigates the formation of protein adducts resulting from cocaine and morphine biotransformation using in vitro assay systems. Human liver microsomal preparations were incubated for 1.5 or 6 h with cocaine or morphine in the presence of thiol-containing trapping agents at 37°C, pH 7.4. Thiols included N-acetylcysteine (NAC), glutathione (GSH), and a synthetic hexapeptide (AcPAACAA). Microsomes were removed by centrifugation and supernatants were subjected to LC-MS/MS analysis for characterization of metabolites and adducts. Isomeric hydroxycocaine adducts from thiol adduction on the azore ring were the major products with all three model thiols. Eight isomers of cocaine-adducted NAC were separated and characterized. While the structural complexity of adducted GSH and model peptide diminished the ability to separate isomers, MS/MS data supported adduct structures analogous to those with NAC. With morphine, two distinct metabolites were identified as the likely species responsible for thiol adduction, the known reactive metabolite morphine and a novel metabolite, morphine quinone methide. Reaction between morphine and NAC formed two isomeric products which underwent secondary reduction to form three additional stereoisomeric products. Likewise, morphine quinone methide formed two structural isomers, although no secondary reduction was noted. Individual structural isomers of morphine adducts with GSH and AcPAACAA could not be isolated; however, morphine-derived adduction products were nevertheless present. Analysis using recombinant cytochrome P450s determined that formation of cocaine adduction products was mediated by CYP1A2, 2C19, and 2D6, while those from morphine were produced by CYP3A4. Results obtained from this study enhance the existing knowledge of illicit drug metabolism and demonstrate novel mechanisms for covalent protein modification by these compounds.
300 Hyperoxia Attenuates Cytochrome CYP1B1 Expression in Human Bronchial Epithelial Cell BEAS-2B: Implications for Oxygen-Mediated Lung Injury.

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Supplemental oxygen, used to treat premature infants with pulmonary insufficiency, contributes to the development of bronchopulmonary dysplasia (BPD) in animal models and infants by mechanisms that are not entirely known. We recently observed that cyp1b1-null mice are less susceptible to hypoxic lung injury, suggesting a pro-oxidant role for CYP1B1. Hyperoxia inhibits the growth of the cells, and β-naphthoflavone (BNF) was reported to protect cells from hyperoxic injury. This study tested the hypotheses: 1. hyperoxia attenuates endogenous and BNF inducible CYP1B1 expression in human lung cell line, BEAS-2B; 2. downregulation of CYP1B1 protects cells from hyperoxic injury while overexpression augments the damage. BEAS-2B cells treated with DMSO (control) or BNF were maintained in room air or hyperoxia for 24, 48, and 72 h. CYP1B1 promoter activity, mRNA and protein expression were evaluated. Cell proliferation, cell viability, apoptotic markers and reactive oxygen species were assessed.

CYP1B1 cells expressed endogenous CYP1B1 protein, which was diminished by about 50% by 24 or 48 h of hyperoxia. BNF induced CYP1B1 mRNA and protein expression. Hyperoxia attenuated endogenous and BNF inducible CYP1B1 protein expression and mRNA expression. Also, hyperoxia attenuated luciferase driven CYP1B1 promoter activity. BNF had minimal improvement in cell viability. Downregulation of CYP1B1 using siRNA improved cell viability in hyperoxia, and overexpression of CYP1B1 was associated with a significant decrease in viability. Our finding that hyperoxia decreases CYP1B1 protein, mRNA and promoter expression suggests transcriptional or post-transcriptional mechanisms. The finding that downregulation of CYP1B1 improves cell viability, while the overexpression decreases cell viability supports the role of CYP1B1 as pro-oxidant in hyperoxic injury. As CYP1B1 appears to contribute to lung injury mediated by hyperoxia, understanding the mechanisms of regulation of CYP1B1 may lead to new strategies to prevent or treat BPD.

301 Metabolism of Deltamethrin (DLM) and Trans-Permethrin (TPM) by Human Hepatic and Intestinal Preparations.


Pyrethroids can be metabolized by both cytochrome P450 (CYP) and carboxylesterase (CES) enzymes. DLM and TPM metabolism was studied using the substrate depletion approach in pooled human hepatic S9, microsomes and cytosol and intestinal S9 fractions. Studies with liver S9 in the presence and absence of NADPH indicated that DLM was metabolised mainly by CES enzymes, whereas TPM was metabolised by both CES and CYP enzymes. Rates of DLM and TPM clearance in liver microsomes were 22.3 and 11.9 ml/min/mg protein, respectively, and in liver cytosol were 2.6 and 5.3 ml/min/mg protein, respectively. The hepatic clearance of both pyrethroids is thus due to both microsomal and cytosolic enzymes. Addition of cofactors for glucuronidation and sulfation did not enhance the metabolism of DLM and TPM by hepatic S9, suggesting that the rate limiting step for hepatic clearance of both pyrethroids is predominantly due to phase I enzymes and not to phase II enzymes. Rates of DLM and TPM clearance by intestinal S9 were 0.6 and 0.8 ml/min/mg protein, respectively. Rates of pyrethroid metabolism were not reduced in the absence of NADPH, suggesting that both compounds are largely metabolised by only intestinal CES enzymes. The metabolic data obtained with human tissue fractions will be used to develop PBPK models for DLM and TPM (Supported by CAPHRA).

302 Biological Impact of a Dysfunctional CYP1/AhR Auto-Regulatory Feedback Loop.

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The toxicity of slowly metabolized AHR agonists (e.g., dioxins) can be explained by their persistent activation of the receptor, whereas transient AHR activation by readily metabolized chemicals leads to toxicity through cytochrome P450 (CYP1)-dependent bioactivation of PAHs to toxic products. Still, CYP1A inhibition has been shown to amplify carcinogenic and teratogenic effects of PAHs, emphasizing the complex relationship between CYP1 induction and toxicity. The endogenous and proposed physiological ligand 6-formylindolo[3,2-b]carbazole (FICZ) has the highest AHR affinity found to date and is an almost perfect substrate for CYP1A, resulting in an efficient auto-regulatory feedback of its actions. The importance of CYP1/AHR feedback regulation to in vivo responses to FICZ is unknown.

We tested the hypothesis that blocking CYP1A expression would result in FICZ becoming toxic in vivo. Studies were performed using zebrarish [zf; Danio rerio] embryos with morpholino knockdown of CYP1A (CYP1A-KD) or AhR2 (AH2-KD). Zf embryos were exposed to vehicle (DMSO) or FICZ (10 or 100nM) starting at 24 h post fertilization (hpf), and morphology was monitored from 54 to 96 hpf. In the CYP1A-KD embryos FICZ caused a dose-dependent increase in the incidence and severity of pericardial edema and circulation failure, and increased lethality. Hatching frequency was reduced and swimbladder inflation abolished. In control-KD and AH2-KD embryos, FICZ (100nM) had no significant morphological effects.

The results show that a functioning CYP1/AHR feedback loop is crucial for regulation of AHR signaling by a potential physiological ligand. Considering the large number of chemicals and drugs known to inhibit CYP1A, we suggest a novel mechanism of toxicity whereby chemicals inhibit the metabolism of FICZ, resulting in prolonged activation of the AHR. [FORMAS grant 2011-963; NIH grants R01ES015912, F32ES017585, and R01ES006272; JSPS Postdoctoral Fellowship for Research Abroad 820]

303 Metabolism and Disposition of 2-Ethylhexyl-p-Methoxycinnamate in Male and Female Harlan Sprague-Dawley Rats and B6C3F1/N Mice After Gavage and Intravenous Administration.

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2-Ethylhexyl-p-methoxycinnamate (EHMC) was nominated to the National Toxicology Program for toxicological evaluation based on its presence as one of the most common active ingredients in sunscreens. Therefore, the current study was undertaken to investigate the metabolism and disposition of [14C]EHMC in male and female Harlan Sprague-Dawley rats and B6C3F1/N mice 24 h or 72 h following gavage and intravenous administration. Intravenous doses to male rats and mice were 8 mg/kg. Gavage doses of 8, 80 or 800 mg/kg to rats were mostly excreted in urine (73-80% in 72 h), with 3-8% of the radioactivity recovered in feces and 1-4% as CO2; volatiles accounted for less than 0.25% of the radioactivity for the 800 mg/kg dose. Radioactive residues in tissues were <1% of the dose. There were no sex or route differences in disposition in rats. In male and female mice administered 8 mg/kg gavage and intravenous doses of EHMC, radioactivity was excreted mostly in urine (57-77% in 72 h), recovery in CO2, and volatiles traps was only 2-4%, respectively, and tissues contained <0.3% of the radioactivity 72 h post dosing without any apparent sex- or route-related differences in disposition. Urinary metabolites following gavage administration of 800 mg/kg EHMC were associated with hydrolysis of the ester and hydroxylation of the ring; no parent EHMC was detected in urine. The metabolites 2-ethylhexanol and 2-ethylhexanoic acid, which are developmental toxicants, were identified by GC-MS in plasma from rats 1 and 2 h following a gavage dose of 800 mg/kg EHMC. These data indicate that oral doses of EHMC are well absorbed, completely metabolized and excreted chiefly in urine with no species or sex difference. [Supported by NIH, N01ES75563]
Immunoochemical Characterization of Xenobiotic-Metabolizing Enzyme Expression in Adult Rat Testis.

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Relatively little is known about the protein expression of xenobioc-metabolizing enzymes, such as cytochrome P450 (CYP) and epoxide hydrolase (EH) in rat testis. These enzymes are expressed in liver and many other organs, and are known to play an important role in the oxidative biotransformation of various endogenous and exogenous compounds. Some xenobioctics such as benzo[a]pyrene are bioactivated to form genotoxic and/or carcinogenic metabolites. Formation of reactive metabolites in the testis could cause severe adverse effects on steroidogenesis and germ cell development. In the present study, we characterized the expression of various xenobioc-metabolizing enzymes in adult rat testis using immunoblot and immunohistochemical analyses. Testicular microsomes prepared from adult male Sprague-Dawley rats were separated using SDS-PAGE and electrophoretically transferred onto membranes and probed with different antibodies. Immunoblot results indicated that CYP1B1, CYP2A1, NADPH-cytochrome P450 reductase, and EH were expressed in testicular microsomes isolated from adult rats. By comparison, CYP1A1, CYP1A2, CYP2B1, CYP2E1, CYP2D1, CYP2D2, CYP2C6, CYP2C7, CYP2C11, CYP2C12, CYP2C13, CYP3A1, CYP3A2, CYP4A1, CYP4A2 and CYP3A4 were not detected in the testicular microsomal samples. In addition, tissue sections were prepared from frozen adult rat testis and probed with antibodies to CYP1B1, CYP2A1, NADPH-cytochrome P450 reductase, and EH. Fluorescent staining indicated that CYP1B1 and CYP2A1 were expressed in interstitial cells, which are comprised mainly of Leydig cells, but not in seminiferous tubules. In contrast, EH and NADPH-cytochrome P450 reductase were expressed in both interstitial cells and in seminiferous tubules. In summary, among the CYP enzymes studied, only CYP1B1 and CYP2A1 were detected in testicular microsomes and appeared to be confined to interstitial cells.

Demographic Differences by Age, BMI, Gender, and Disease States of Phase I and Phase II Enzyme Activities in Cryopreserved Human Hepatocytes.


Human hepatocytes are a key in vitro reagent for making predictions of in vivo drug metabolism, interactions and intrinsic clearance in drug discovery and development. However, inter-individual differences in drug metabolizing enzyme activities complicate pharmacokinetics, leading to varying efficacy and drug-drug interactions. To delineate the potential influences, we have reviewed phase I (CYP1A2, CYP2E1, CYP2B6, CYP2C9: tolbutamide, CYP2C19: mephenytoin, CYP2C9: phenacetin, CYP2C19: tolbutamide, 7-ethoxycoumarin (ECOD), 2C9: tolbutamide, 2C19: mephenytoin, 2C8: chlorzoxazone, 3A4: testosterone and other substrates) and phase II (N-acetyltransferase 1, 2, 3-glucuronidase, 6-glucuronidase) drug metabolism and drug interactions. As BMI increased, ECOD, CYP1A2 and CYP2C19 decreased be associated with multiple enzyme substrate 7-ethoxycoumarin (ECOD).

Metabolism of Benzo(a)pyrene (BAP) and Fluoranthene (FLA) in Gastrointestinal Tract Subcellular Fractions of the APCMin Mouse.


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The objective of the present study was to investigate whether subcellular fractions (nuclear, cytosolic, mitochondrial, and microsomal) from the gastrointestinal (GI) tract of a colon cancer mouse model were capable of metabolizing BAP, a combustion byproduct, which is released into the environment from automobile exhausts, cigarette smoke, and industrial emissions. A significant intake of BaP is also expected in people who consume barbecued foods, and diet rich in saturated fat. In this study, subcellular fractions (SCF) were isolated by differential centrifugation from a tumor-bearing ApcMin mice induced by subchronic doses of 500ug/kg BaP and incubated with either BaP or FLA (3mM each) alone or in combination and appropriate control groups. Subsequent to incubation, samples were extracted with ethyl acetate and analyzed for BaP and FLA metabolites by reverse-phase HPLC with fluorescence detection. The SCF from tumors tissues metabolized BaP to a greater extent than those from the non-tumor tissues. The rate of BaP metabolism (pmol of metabolite/min/mg protein) was found to be when more fractions from BaP-pretreated mice were exposed to BaP alone. The SCF from BaP-pre-exposed mice generated greater proportion of BaP, 7,8-diol, BaP 3,6- and 6,12-diones compared to other experimental groups. Furthermore, SCF from BaP-pretreated mice produced greater proportion of FLA 2, 3-diol, and 2, 3 D FLA when fractions were incubated with FLA alone or a combination of BaP and FLA. Our studies revealed that the tumor SCF were potent to metabolize BaP and FLA either individually or as a binary mixture. The metabolism of BaP and FLA as a consequence of prior or simultaneous exposure to BaP may influence the growth of tumors. Our findings are of relevance to long-term dietary intake of these toxicants and the consequent acceleration of the GI tract carcinogenesis process in humans (supported by S05CA142445-02, 1F31ES017391-01, 1F31ES019432-01A1, 5R25GM059994-11 and SREB grants).

Studies of Sterylene, Sterylene Oxide and 4-Hydroxystereane Toxicity in CYP2F2 Knockout and CYP2F1 Humanized Mouse Support Lack of Human Relevance for Mouse Lung Tumors.

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Sterylene (S) is lung tumorigenic in mouse but not in rats. In previous mode of action (MOA) studies, S and its alkene-oxidized metabolite styrene oxide (SO) were not lung toxic in CYP2F2(-/-) [knockout] mice, indicating S-induced mouse lung tumors are mediated through mouse-specific CYP2F2-generated cytotoxic ring-oxidized metabolite(s) producing repeated localized cytotoxicity in Clara cells and associated cumulative cell proliferation in lung bronchioles. This conclusion is consistent with the observation that 4-hydroxystereane (4HS) is toxic to Clara cells in culture at doses lower than S or SO. We hypothesized that CYP2F2 MOA was assessed by insertion of a human CYP2F1,2A13,2B6 transgene into CYP2F2(-/-) mice; CYP2F1 expression and activity were confirmed in the transgenic (TG) mice. No evidence of cytotoxicity or increased cell proliferation (BrDU labeling) was seen in TG mice treated with either S or SO (200 mg/kg/d for 5 days), while cytotoxicity was apparent and BrDU labeling was increased 10-fold in wild-type (WT) mice. Consistent with the hypothesis, 4HS (60 or 105 mg/kg ip for 5 days) increased BrDU labeling 5-10 fold in WT mice, and was attenuated to less than a 3 fold increase in KO mice and a 2-4 fold increase in TG mice. The limited response of 4HS in both KO and TG mice suggests that direct administration of high doses of 4HS are either intrinsically lung toxic or are capable of limited metabolism by either CYP2F1 and/or other lung CYPs in KO and TG mice to lung toxic metabolite(s). Regardless of the MOA of the limited 4HS toxicity in KO or TG mice, these findings indicate that the CYP2F2-mediated tumorigenic MOA in WT mice is not operative for S, SO, or for 4HS putatively derived from metabolism of S by CYP2F2 in humans, and thus S-induced mouse lung tumors are unlikely to be relevant to human risk.

Sponsor: Sterylene Information and Research Center

Metabolism and Disposition of Bisphenol AF in Male and Female Harlan Sprague-Dawley Rats following Gavage and Intravenous Administration.

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Bisphenol AF (BPAF) is used as a cross linking agent in polymers. BPAF was nominated to the National Toxicology Program for toxicological evaluation based on its moderate production levels, structural similarity to bisphenol A, and lack of adequate toxicity data. The current study was undertaken to investigate the metabolism and disposition of [14C]BPAF in male and female Harlan Sprague Dawley (HSO) rats. Following gavage administration of 3.4, 34 or 340 mg/kg to male rats, the administered dose was mostly excreted in feces (73-80%) with < 6% of the dose recovered in urine and cage rinse at 72 h. Radioactivity in tissues was 0.2-1.5% of the dose 72 h post dosing compared to 7% at 24 h following a dose of 34 mg/kg.
Urinary excretion at 72 h post administration was higher in females (-15%) compared to males (-4%) following a 34 mg/kg dose. Distribution of an intravenous dose of 34 mg/kg was similar to that following gavage, but with less excretion in urine in both males (0.62% for intravenous vs. 4.32% for gavage) and in females (7% for intravenous vs. 15% for gavage). About 52% of a 340 mg/kg gavage dose was excreted in bile by 24 h, indicating that high excretion in feces is not due mostly to gastrointestinal absorption. BPA metabolite glucuronide (major metabolite), diglucuronide, glucuronide-sulfate, and sulfate were identified in bile using LC/MS/MS. This study demonstrated that BPA is well absorbed following gavage administration in rats, metabolized by glucuronidation and sulfation, and excreted mainly in feces. [Supported by NIH, N01ES75563].

PS 309 Absorption, Distribution, Metabolism, and Excretion Studies of n-Butylbenzenesulfonamide in Harlan Sprague-Dawley Rats Following Gavage Administration.

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n-Butylbenzenesulfonamide (NBBS) is used as a plasticizer and an antifungal agent. There is high potential for human exposure to NBBS due to its likely occurrence in drinking water and leaching from NBBS-containing products. The limited toxicity data in rodents suggests that NBBS can cause toxicity to the hematopoietic, nervous, and male reproductive systems. The present studies were conducted to investigate the clearance of NBBS in male Harlan Sprague Dawley (HSD) rat and B6C3F1/N mouse hepatocytes in vitro and metabolism and disposition of NBBS following gavage administration in HSD rats in vivo. The half-lives of disappearance of NBBS (1 μM) in rat and mouse hepatocytes were 136 ± 24 and 320 ± 41 min, respectively. Following gavage administration of ring-labeled [14C]NBBS to male HSD rats at 2 and 200 mg/kg and sacrificed at 72h, NBBS was excreted primarily in urine (70-76%) with feces accounting for about 12% of the administered dose. Retention in tissues was 5.7-7% at 72 h, with no tissue exhibiting high concentrations of radioactivity relative to blood. Profiling of urine showed presence of numerous metabolites; however, the parent NBBS was not observed. Some of the polar metabolites were diminished upon treatment with β-glucuronidase or acetylase, indicating glucuronides and metabolites that minimize characterize, which are not uncommon in such processes. In conclusion, NBBS is well absorbed following gavage administration and metabolized to products including glucuronides and mercapturates. [Supported by NIH, N01ES75563].

PS 310 Biotransformation of BDE-47 to Potentially Toxic Metabolites Is Predominantly Mediated by Human CYP2B6: Implications for Interindividual Variability in Metabolism and Disposition of BDEs.

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Recent studies suggest that bioactivation by oxidative metabolism may add considerably to the neurotoxic potential of polynuclear aromatic hydrocarbons (PAHs) and their bio-active metabolites. We investigated the potential for oxidative metabolism of BDE-47, the most abundant PAH detected in human urine, to produce toxic metabolites. Seven human cytochrome P450s (CYPs), and to identify P450s that are active in the oxidative metabolism of BDE-47. Human CYPs (CYP1A1, CYP1A2, 1B1, 1A6, 2B6, 2C8, 2C9, 2C19, 2D6, 2E1, 3A4) were incubated with BDE-47 (20 μM) and metabolites were measured and characterized using GC/MS/MS. For kinetic studies, CYP2B6 and pooled HLMs were incubated with BDE-47 (0-60 μM). CYP2B6 was the predominant CYP capable of forming six different OH-BDEs, including 3-OH-BDE-47, 5-OH-BDE-47, 6-OH-BDE-47, 2′-OH-BDE-66, 4-OH-BDE-42, and 4′-OH-BDE-49. GC/MS analysis also revealed formation of novel metabolites, di-OH-BDE-47 and di-OH-dioxin. Kinetic studies of BDE-47 metabolism by CYP2B6 and pooled HLMs found Km values ranging from 4.9-9 μM and 7.8-13 μM, respectively. CYP2B6-specific metabolism may contribute to interindividual variability in the body burden of PBDEs and the formation of potentially toxic metabolites. (NIEHS, grant # ES021554)
present study is to evaluate the potential protective effects of EGCG on benz[a]pyrene (BaP)-induced cytotoxicity and DNA damage in BEAS-2B, a human normal lung epithelial cell. BEAS-2B cells were treated with vehicle control (0.1% DMSO), BaP or BaP+EGCG for 24 hours. The cytotoxicity, cell cycle, benzo[a]pyrene diol epoxidation (BPDE)-DNA adducts, and mRNA expression levels of cytochrome P450 (CYPs) were determined by MTT assay, flow cytometry, high performance liquid chromatography (HPLC), and quantitative real-time PCR (qRT-PCR), respectively. BaP induced cell growth inhibition in a dose-dependent manner; while EGCG dose- dependently reversed this inhibition (P<0.05). The flow cytometry analysis showed that BaP caused significant G2/M phase arrest compared to controls, however, increased S phase and decreased G2/M phase were observed in cells co-treated with BaP and EGCG compared to BaP group (P<0.05). BEAS-2B cells exposed to BaP had a significant induction of BPDE-DNA adducts when compared with controls (P<0.01). Moreover, these adducts were diminished significantly by EGCG treatment with an 80% reduction. CYP1A1 and CYP1B1 expression levels analyzed by qRT-PCR dramatically increased after BaP exposure compared to controls (CYP1A1: 130.2-folds; CYP1B1: 6.0-folds; P<0.001). EGCG significantly reduced BaP-induced CYP1A1 and CYP1B1 expression (CYP1A1: 1.6-folds; CYP1B1: 1.2-folds; BaP vs. BaP+EGCG, P<0.05). On the other hand, CYP1A2 and CYP3A4 did not show any changes among the control, BaP-treated, and BaP and EGCG co-treated groups. In summary, BaP-induced adverse effects could be prevented by EGCG, suggesting a possible chemopreventive role for this natural polyphenol against the development of lung cancer.

314 17β-Estradiol Benzoate and Bisphenol A Suppress Xenobiotic-Metabolizing Enzyme Expression in Adult Rat Testis.

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Previous studies showed that 17β-estradiol benzoate (EB) (at 4 μmol/kg) decreased testicular expression of cytochrome P450 1B1 (CYP1B1) in adult male rats. Bisphenol A (BPA) is an endocrine disrupting chemical that has been reported to exert estrogenic activity in vitro and in vivo. In the present study, we investigated the effect of treatment with EB and BPA at varying dosages on the expression of CYP and other xenobiotic-metabolizing enzymes in rat testis. In Experiment 1, five groups of adult male Sprague-Dawley rats (n = 4 per group, except for control, n = 5) were injected sc with EB (0.004, 0.04, 0.4 or 4 μmol/kg) or vehicle (propylene glycol, 1ml/kg), once daily for 14 days. Immunoblot analysis indicated that treatment with EB at 0.004, 0.04, 0.4 or 4 μmol/kg decreased testicular CYP1B1 expression (by 15, 56, 70 and 80 %), CYP2A1 (by 48, 79, 97 %) and CYP17A1 (by 15, 44, 94 and 98 %) protein levels, respectively, when compared with vehicle-treated rats. The constitutive expression of EH and NADPH-cytochrome P450 reductase was not affected by EB at 0.004 μmol/kg, but EB (by 56, 70 and 80 %) and NADPH-cytochrome P450 reductase (by 53, 58 and 48 %) protein levels were decreased following EB treatment at 0.04, 0.4 and 4 μmol/kg, respectively. In Experiment 2, adult male Sprague-Dawley rats (n = 4 per group, except for control, n = 5) were injected sc with propylene glycol (1ml/kg) or BPA at 400, 800 or 1600 μmol/kg, once daily for 14 days. Treatment with BPA at 400, 800 or 1600 μmol/kg decreased CYP1B1 (by 51, 87 and 89%), CYP2A1 (by 79, 92 and 92%), EH (by 50, 67 and 67%) and NADPH-cytochrome P450 reductase (by 43, 67 and 67%) protein levels, respectively, when compared with saline- or vehicle-treated rats. CYP1A1 protein levels were decreased by 49% by BPA at 400 μmol/kg dose and were not detectable at other BPA doses. In conclusion, testicular expression of CYP1B1, CYP2A1, CYP17A1, EH and NADPH-cytochrome P450 reductase was down-regulated by exogenous estradiol and BPA, at higher dosages, produced a similar effect.

315 Acute Lead Exposure Induces Cardiotoxicity In Vitro and In Vivo Rat Model through the Cytochrome P450 1A1 Signaling Pathway.

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Activity-based protein profiling (ABPP) has recently emerged as a post-genomic technology for characterizing functional proteins in complex biological systems. This approach applies chemical activity-based probes that monitor the functional activity of enzymes under physiological conditions, thereby providing high-content proteomic information beyond the reach of standard gene- and protein-expression profiling. We have previously developed a set of activity-based probes that display broad coverage across mouse and human phase I and II metabolizing enzymes. Each probe contains three moieties: (1) a reactive group that forms a covalent bond to a functional P450 through a mechanism-based reaction, (2) a binding group that targets a probe towards P450s, and (3) an alkyne (C2) handle to exploit the bio-compatible click chemistry reaction for attachment of an enrichment moiety or fluorophore reporter. In this study, we measured enzyme activity at key life stages in mouse, including fetal development, normal adulthood, and pregnancy, as well as in human fetal tissues. Additionally, we compared the effects of exposure to the transplacentral carcinogenic polycyclic aromatic hydrocarbon (PAH) dibenzo[def,p]chrysene (DBC) on enzyme activity in mouse. We have determined that the activity of most hepatic phase I enzymes associated with PAH metabolism (e.g., CYP 1A1, CYP2E1, epoxide hydrolase) was reduced by 2-10 fold during pregnancy in mice. By incorporating these reductions in enzyme activity

316 Reciprocal Roles of Cytochromes P4501a1 and 1a2 in Lung Carcinogenesis Mediated by Polyyclic Aromatic Hydrocarbons (PAHs) in Mice.

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Humans are constantly exposed to polycyclic aromatic hydrocarbons (PAHs). Cytochrome P4501a1 (CYP1A1) enzymes play important roles in the activation of PAHs such as 3-methylcholanthrene (MC) to DNA-binding metabolites, which in turn mediate carcinogenesis in target organs such as lung. In this study, we tested the hypothesis that CYP1A1 and 1A2 have reciprocal roles in PAH-mediated tumorigenesis. Eight week-old female wild type (WT) (A/J) mice or mice lacking the gene for CYP1A1 or CYP1A2 on the A/J background were treated with a single dose of MC (40 μmol/kg), or vehicle (corn oil), and liver and lung tumors were studied after 28 weeks. While 100% of WT or Cyp1a2-null mice exposed to MC showed lung tumors after 28 weeks, about 80% of the Cyp1a2-null mice showed lung tumors. However, there were striking differences in the Cyp1a1-null and Cyp1a2-null mice in regard to lung tumor multiplicities. The WT mice treated with MC had about 15 lung tumors/animal. On the other hand, the Cyp1a2-null mice displayed about 40 lung tumors/animal. In Cyp1a1-null mice, about 2-3 tumors/mouse. DNA adduct studies at early time points (8 days) showed increased MC-DNA adducts in lungs of Cyp1a2-null mice compared to WT mice, and decreased adduct formation in the Cyp1a1-null mice, supporting the hypothesis that DNA adducts are early biomarkers of PAH-mediated carcinogenesis. Overall, our results suggest that CYP1A1 contributes to the formation of tumorigenesis by PAHs, while CYP1A2 protects against carcinogenesis mediated by PAHs, presumably through it role in PAH detoxification. In conclusion, our results strongly suggest that CYP1A1 and 1A2 could be novel candidates for cancer prevention and therapy though either inhibition of CYP1A1 or induction of CYP1A2. (Supported by NIH grant E509132.)

317 Activity-Based Protein Profiling of Metabolizing Enzyme Ontogeny and Response to PAH Exposure in Rodents and Humans.

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Activity-based protein profiling (ABPP) has recently emerged as a post-genomic technology for characterizing functional proteins in complex biological systems. This approach applies chemical activity-based probes that monitor the functional activity of enzymes under physiological conditions, thereby providing high-content proteomic information beyond the reach of standard gene- and protein-expression profiling. We have previously developed a set of activity-based probes that display broad coverage across mouse and human phase I and II metabolizing enzymes. Each probe contains three moieties: (1) a reactive group that forms a covalent bond to a functional P450 through a mechanism-based reaction, (2) a binding group that targets a probe towards P450s, and (3) an alkyne (C2) handle to exploit the bio-compatible click chemistry reaction for attachment of an enrichment moiety or fluorescent reporter. In this study, we measured enzyme activity at key life stages in mouse, including fetal development, normal adulthood, and pregnancy, as well as in human fetal tissues. Additionally, we compared the effects of exposure to the transplacentral carcinogenic polycyclic aromatic hydrocarbon (PAH) dibenzo[def,p]chrysene (DBC) on enzyme activity in mouse. We have determined that the activity of most hepatic phase I enzymes associated with PAH metabolism (e.g., CYP 1A1, CYP2E1, epoxide hydrolase) was reduced by 2-10 fold during pregnancy in mice. By incorporating these reductions in enzyme activity
into a physiologically based pharmacokinetic (PBPK) model, along with normal changes in anatomy and physiology, we were able to describe the elevated concentrations of DBC in blood and tissues of pregnant mice versus naïve mice, following equivalent exposures. We additionally report, for the first time, developmentally driven changes in enzyme activity in both mice and humans. Supported by Award Number P42 ES016465 from the NIEHS, and DOE Laboratory Directed R&D Project 90001.

318 Eliciting the Catalytic Mechanisms of the Novel Sterylene Detoxification System in Pseudomonas Bacteria.


As the world becomes more polluted with styrene-based polymers, it is essential to consider how the 28 million tons of Styrofoam and commercial plastics produced in the United States impact human health. Sterylene and its metabolites act as biological membrane disruptors and intracellularly where they are transformed by cytochrome p450 isozymes to styrene oxide and vinylphenol, which have biological activity as pulmonary and hepatic poisons. This toxic vinyl benzene is a danger to living systems and for this reason we focused our research on the styrene detoxification pathway in Pseudomonas bacteria. The first enzyme, styrene monoxygenase (SMO), a two component flavoenzyme with an NADH specific reductase, SMOB, and a FAD specific epoxidase, SMOA, catalyzes the epoxidation of styrene to yield styrene oxide. Here, SMOB binds NADH and oxidized FAD as substrates and catalyzes the reduction of FAD by a hydride-transfer mechanism. Many details of the SMOB catalyzed flavin reduction and transfer still need to be explained. This research evaluates the catalytic mechanism of N-terminally histidine-tagged styrene monoxygenase reductase (N-SMOB) from Pseudomonas putida S12 bacteria. Over expression of N-SMOB in E.coli BL21(DE3) cells produces high amounts of the enzyme, which we purified using nickel affinity chromatography. A Spectromax190 microplate reader was used to measure the rate at which the hydride ion is transferred from NADH to FAD at 340nm. Previous data showed the native SMOB enzyme follows a sequential mechanism under identical conditions. However, preliminary data of N-SMOB at 10 °C suggests that a double displacement reaction with NADH as the leading substrate could be prevalent mechanism. These studies of N-SMOB at 10°C provided estimates of the Km of NADH, Km of FAD and Vmax of N-SMOB at 5.0 uM, 3.7 uM, and 38.0 uMs⁻¹, respectively. Through the use of high resolution kinetic analysis at 30°C, we confirm that the double displacement mechanism is the preferred reaction for N-SMOB and these findings will be used to elucidate the flavin-transfer reaction.

319 Impact of Liver-Specific Loss of Cytochrome P450 Reductase on the Expression and Function of Intestinal P450 Enzymes.

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Tissue-specific deletion of the cytochrome P450 reductase (CPR) gene in either liver or intestine leads to upregulation of many P450 genes in the tissue with the Cpr deletion. The aim of this study was to test whether a loss of hepatic CPR would also lead to upregulation of P450 expression in the small intestine (SI). We found that in the liver-specific Cpr-null (LCN) mouse, SI expression of CYP2B, 2C and 3A proteins was increased, by 2- to 5-fold, relative to that in wild-type (WT) mice. This increase was accompanied by increases in rates of SI microsomal metabolism oflovastatin (LVS), a CYP3A substrate (by 2.1-fold). The overall impact of the loss of hepatic CPR/P450 function and the increase in SI P450 expression on systemic clearance of orally administered LVS was dependent on LVS dose: the rates of clearance were increased at an LVS dose of 5 mg/kg, unchanged at 25 mg/kg, but decreased at 50 mg/kg, in the LCN mice, compared to WT mice. Thus, we show for the first time that hepatic CPR/P450 deficiency leads to compensatory increases in SI P450 expression and capacity of first-pass metabolism of oral drugs; this finding may aid in the prediction of drug exposure in patients with compromised hepatic P450 function. We also found that SI FGF15 and IBABP mRNA levels were significantly lower (by 15- and 5-fold, respectively) in LCN than in WT mice, while the levels of PXR and FXR mRNAs were unchanged. Furthermore, treatment of mice with FGF19 (the human counterpart of mouse FGF15) abolished the difference between WT and LCN mice in SI CYP3A levels. Thus, SI FGF15 may play a previously unrecognized, direct role in the regulation of SI P450 expression. Overall, our results not only provide the basis for further mechanistic studies of physiological regulation of SI P450 expression, but will also help with data interpretation for numerous studies utilizing the LCN mouse model to determine roles of hepatic P450s in the disposition of orally administered drugs.

320 Organ-Specific Ugt1 Locus Profiling in Defining the Toxic Response Towards Irinotecan Anticancer Drug Therapy.

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Irinotecan (CPT-11) has been used as a first line drug in the treatment of colorectal cancer. However, its efficacy and safety is compromised because of severe late onset diarrhea, a result of enterocyte toxicity from the active metabolite SN-38. SN-38 is inactivated primarily by hepatic UGT1A1 catalyzed glucuronidation to form SN-38 glucuronide, which is excreted via the biliary ducts into the gastrointestinal (GI) tract, where it serves as a substrate for bacterial β-glucuronidase. Free SN-38 is then re-absorbed by entry through the GI tract. Since an abundance of the UGT1 proteins are rich in the GI tract, it was important to examine the association between SN-38 glucuronidation and the pending intestinal tissue damage resulting from CPT-11 therapy. To carry out these experiments, we have generated mouse models targeting deletion of the Ugt1 locus specifically in liver (Ugt1Δliver) and the intestines (Ugt1Δintestine). Wild type (Ugt1ΔWT), Ugt1Δliver, and Ugt1Δintestine adult male mice were treated by the intraperitoneal route with CPT-11 daily for four constitutive days. At a daily dose of 75 mg/kg, survival curves of the Ugt1Δliver mice showed a 50% lethality rate, comparable to the LD50 values of CPT-11 treated Ugt1ΔWT mice. Deletion of the Ugt1 locus in hepatic tissue had no impact on liver or GI toxicity when compared to wild type mice. Alternatively, Ugt1Δintestine mice were highly susceptible to CPT-11-induced diarrhea, developing severe ileoclitus. At a CPT-11 dose of 25 mg/kg, bloody diarrhea was observed in all Ugt1ΔWT mice and was associated with 100% lethality, while no diarrhea or lethality was observed in Ugt1Δliver or Ugt1ΔWT mice at that dose. Thus, intestinal expression of the UGT1A proteins is critical towards the detoxification of SN-38. Regulation of the intestinal UGT1A1 gene may serve as a target for improving the therapeutic index and efficacy associated with CPT-11 treatment. (Supported by USPHS grants ES010337 and CA171008)


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Cytochrome P450 (CYP) 3A4 is one of the major isozymes of enzymes that catalyze Phase I oxidation a wide variety of endogenous and exogenous compounds (i.e., broad substrate specificity) including environmental xenobiotics like chiral pesticides. In particular, several chiral pesticides are well-known to undergo differential kinetics within the homochiral environment of biological systems. The current ability to model such metabolic processes within physiologically-based pharmacokinetic model from an exposure-dose perspective is limited by specific chiral information coupled together with accurate kinetic data. With available in-house stereospecific (i.e., individual stereoisomer) in vitro data, we have utilized a ligand-based approach to develop a prototype CYP3A4 pharmacophore model for triazole fungicides based on the observed stereoselective CYP3A4 clearance. The developed model was further used to discriminate between single isomeric configurations thereby enriching the dataset and providing further criteria on putative ligand-receptor interactions. We also utilized a combined pharmacophore and docking approach within the CYP3A4 binding cavity (2V0M FDBID) to provide estimates on putative ligand stereoselective rate constants for a test set of triazole compounds. Observations and comparisons between high and low binding affinity poses as well as homolytic dissociation bond energy estimates at purported sites of metabolism indicate that this method may be helpful for selecting catalytically competent ligand substrates for CYP3A4 and thereby reducing the uncertainty within exposure-based pharmacokinetic models of chiral metabolism. Although this work was reviewed by EPA and approved for publication, it may not necessarily reflect official Agency policy.

322 Chlorpyrifos Affects Specific Types of Zebrafish Larval Behavior If Administered during Distinct Developmental Time Periods.

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Pesticides are widely used in agriculture and are found ubiquitously in the environment. While adults have enzymes that are able to break down pesticides, developing embryos lack the necessary enzymes for toxicant removal. Low doses of organophosphate pesticides during early embryonic development in animal models...
and in humans have been documented to affect brain development and behavior. It is important to determine at which stages of development embryos are most affected by the exposure to organophosphate pesticides. Using zebrafish as a model system for pesticide exposure is advantageous because embryos can be exposed to organophosphates immediately after fertilization and large numbers of larvae can be used for high-throughput behavioral analysis. Behaviors such as swim speed, preference for edge of a well, and avoidance of a moving stimulus can be quickly obtained. In the present study, employing a high-throughput assay unique to our lab, we show that exposure to low doses of a widely used organophosphate, chlorpyrifos, affects discrete behaviors if administered during specific developmental time periods. The results indicate that even low levels of chlorpyrifos have the potential to impact behavior when administered during critical periods of development. These behavioral abnormalities are emergent from pesticide exposure during different critical points. The results of the present study have the potential to affect food consumption guidelines, especially in pregnant women. Future studies will include larval zebrafish confocal brain imaging to detect neural patterning changes after chlorpyrifos exposure.

### 323 The AhR Pathway and Aromatic Hydrocarbon-Mediated Teratogenicity in the Atlantic Killifish.

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Exposure of developing fish to polycyclic aromatic hydrocarbons (PAHs) and halogenated aromatic hydrocarbons (HAHs) results in a suite of defects including cardiac malformation, pericardial and yolk sac edema, craniofacial defects and hemorrhaging. Several populations of Atlantic killifish (Fundulus heteroclitus) on the Atlantic coast of the United States are resistant to the developmental toxicity caused by PAHs and HAHs; this resistant phenotype displays strong down-regulation of the aryl hydrocarbon receptor (AhR) pathway. The AhR is known to mediate many toxic responses to PAHs and HAHs in vertebrates. A single AhR has been identified in mammals, but killifish and other fish species have multiple AhRs and their roles in contaminant response and development are not clear. In this study, translation-blocking and splice-junction morpholino gene knockdown was used to determine the roles of AhR1 and AhR2, as well as CYP1A, in mediating cardiac teratogenesis induced by β-naphthoflavone (BNF), benzo[k]fluoranthene (BkF), and 3, 3′, 4, 4′, 5-pentachlorobiphenyl (PCB-126). Here we report the AhR2 and not AhR1 knockdown resulted in rescue of cardiac teratogenicity induced by BNF, BkF, and PCB-126 in laboratory-reared offspring of non-adapted parents from unpolluted sites. Knockdown of CYP1A enhanced the toxicity of PAHs but not PCB126. Furthermore, knockdown of AhR2 partially rescued the severe cardiac teratogenicity caused by extracts of sediments from the Atlantic Wood (AW) Superfund site (Elizabeth River, VA, USA), a site heavily contaminated with a complex mixture of creosote-derived PAHs (and home to a resistant population of killi-fish). These data demonstrate that AhR2 is an important mediator of cardiac teratogenesis caused by multiple aryl hydrocarbons in killifish and suggest that suppression of the AhR pathway through developmental modulation of AhR2 or other mechanisms may play a protective role against PAH developmental toxicity.

### 324 Characterization of g101 Gene Expression during Development and Alterations Induced by Atrazine Exposure in Zebrafish.

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Atrazine is a commonly used herbicide that is an endocrine disruptor and a suspected carcinogen. Although atrazine was recently banned by the European Union for widespread contamination risks in potable water supplies, this herbicide is still commonly used in the United States with a current maximum contaminant level (MCL) of 3 ppb. The health risks associated with this MCL are currently being re-viewed by the Environmental Protection Agency, but the mechanisms of atrazine toxicity are not well defined. In this study, we are using [1] global gene expression analysis to identify altered genes, [2] in situ hybridization to qualitatively analyze toxic responses, and [3] quantitative PCR (qPCR) to assess gene expression levels of specific targets throughout development in the zebrafish model system. In a previous study completed in our laboratory, zebrafish were exposed to 0.3, 3, or 30 ppb atrazine or a control treatment throughout development (-72 hours post fertilization [hpf]). This analysis showed that expression alterations were enriched with genes associated with neuroendocrine development and function, cell cycle regulation, and carcinogenesis. From this list of genes, glyc oxylyase 1 (GLO1) was targeted for further study. GLO1 is part of the glyoxylate system which converts methylglyoxal to S-D-lactoylgluthathione. Uprogelation of GLO1 is linked to cell proliferation and is associated with various cancers in humans. To further our understanding of this genetic target, expression was analyzed at five developmental time points: 24, 36, 48, 60, and 72 hpf. In situ hybridization and qPCR were coupled to determine spatial and quantitative gene expression of g101 throughout development under normal conditions and after atrazine treat-ment. This data is furthering our understanding of g101 expression during development and alterations induced by atrazine exposure.


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Birds show a strikingly large species variation in susceptibility to developmental toxicity of dioxins and coplanar polychlorinated biphenyls (PCBs). The chicken is the most sensitive to these compounds among avian species, while e.g. the Japanese quail is much more resistant. Such differences are proposed to depend on binding affinity of dioxin to the aryl hydrocarbon receptor (AhR), i.e., the degree of sensi-tivity is linked to variation in a few amino acids in the ligand binding domain of avian AhRs. The proposed endogenous AhR ligand 6-formylindolo[3,2-b]carbazole (FICZ) has high AhR binding affinity and causes strong transient induction of CYP1 mRNA in human cells and zebrafish embryos. In Xenopus tropicalis tadpoles FICZ is almost as potent an inducer of CYP1A as in zebrafish embryos. However, Xenopus tadpoles are much less sensitive to PCB126-induced toxicity than zebrafish embryos. The goal of the present study was to determine mRNA expression responses and other effects of FICZ in developing birds. Peanut oil-lecithin emulsions with or without FICZ were injected into the yolks of day-4 chicken embryos (0, 2, 20, or 200 μg FICZ/kg egg) and day-3 quail embryos (0, 2, or 200 μg FICZ/kg egg). Twenty four hours post-injection CYP1A4 and CYP1A5 showed dose-dependent induction by FICZ in both chicken and quail, and the degree of induction was similar in the two species. The CYP1B1 level in both species and the CYP1C1 level in chicken were unaffected by FICZ exposure. In chicken embryos exposed to 200 μg FICZ/kg the CYP1A levels remained induced in liver and thymus 13 days post-injection (day 17). Furthermore, liver lesions were observed in 48% of these animals, suggesting that FICZ is toxic at high doses. Our results suggest that while there is a large species variation in sensitivity to dioxin-like compounds the strong AhR activation by FICZ is evolutionarily conserved, indicat-ing an important physiological function. Funding: The Swedish Research Council Formas and Carl Tryggers Stiftelse.

### 326 Aryl Phosphate Esters within a Major PentaBDE Replacement Induce Cardiotoxicity in Developing Zebrafish Embryos: Potential Role of the Aryl Hydrocarbon Receptor.

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PentaBDE (550) (FM550) is an additive flame retardant formulation of brominated and aryl phosphate ester (APE) components introduced in 2004 as a major replace-ment for the commercial polybrominated diphenyl ether mixture (known as PentaBDE) used primarily in polyurethane foam. Due to rapid adoption, certain FM550 components have been detected at elevated concentrations within indoor environments. However, little is known about the potential effects of FM550-based ingredients during early vertebrate development. Therefore, we first screened the developmental toxicity of each FM550 component using zebrafish as an animal model. Based on these initial screening assays, we found that exposure to triphenyl phosphate (TPP) or mono-substituted isopropyli triaryl phosphate (Mono-ITP) – two APEs comprising almost 50% of FM550 – resulted in targeted effects on cardiac looping and function during embryogenesis. As these cardiac abnormalities resembled aryl hydrocarbon receptor (AhR) agonist-induced phenotypes, we then exposed developing embryos to TPP or Mono-ITP in the presence or absence of a selective AhR antagonist (CH223191) or AhR2-specific morpholino. Based on these studies, we found that CH223191 blocked the cardiotoxic effects of Mono-ITP but not TPP, while AhR2 knockdown failed to block the cardiotoxic effects of both components. Moreover, using a cell-based human AhR reporter assay, we found that Mono-ITP (but not TPP) exposure resulted in a significant in-crease in human AhR-driven luciferase activity at similar nominal concentrations as a potent reference AhR agonist (β-Naphthoflavone). Overall, our findings sug-gest that two major APE components of FM550 induce severe cardiac abnormalities during early vertebrate development, raising questions about the potential health risks of these APEs resulting from indoor exposure.
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Cytorchrome P450 Gene Transcripts in Early Development of Zebrafish Danio rerio.


Understanding the roles of cytochromes P450 (CYPs) in zebrafish is important to the use of this non-mammalian model in toxicological, pharmacological and carcinogenesis research. In this study, we determined whether maternally derived transcriptions for many CYP genes are present in zebrafish oocytes, and how levels change up to the mid-blastula transition (MBT) (3 hours post fertilization, hpf), focusing on genes involved in xenobiotic and endobiotic metabolism. The maternal contribution to transcript abundance in the eggs varied greatly among CYP genes examined. CYP2P1 showed the highest levels in the oocytes, followed by CYP2O and CYP1A. The transcript levels of all CYPs examined were similar between unfertilized eggs and fertilized eggs at 2 cell stage prior to the first division. Many of CYPs including CYP1B1, 1C1, 1C2, 2P6, 2R1, 2A4, 11C1, 17A2, and 26A1 showed significant increases in their transcript levels at 3 hpf (1,000 cells). In contrast, CYP1A and the steroidogenic CYPs 11A1, 17A1, and 19A1, showed constant levels of transcript from egg to 3 hpf. Other CYPs, including CYP2V1, 2A3, 3C1, and CYP51 showed an increase trend at 2 hpf (64 cells), well before the MBT. We also examined the effect of exposure to an ary1 hydrocarbon receptor agonist, 3,3',4,4',5-pentachlorophenol (PCB126) on transcript levels of CYP1 and selected other genes in oocytes. Exposure of females to waterborne PCB126 caused increases in the transcript levels of CYP1A, 2A3, 3C1, 11A1, and CYP51 genes in eggs as compared to those from vehicle-exposed females. The results reveal maternal transcript deposition of a suite of CYP genes in the egg, and also show that temporal patterns of CYP transcript levels during early development from egg to the MBT differ substantially among different CYP genes. Maternal exposure to chemical was shown to cause change in the transcript deposition of some CYPs in the eggs. [Support: JSPS Postdoctoral Fellowships for Research Abroad no. 820 (A.K.), JSPS KAKENHI grants 20686037 and 20150037 (K.M.); NIH Superfund Research Program grant P42ES00738 (J.S.)]

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Benzo[a]pyrene Exposure Effects on Reproductive Success, Development, and Transcriptome in Zebrafish.

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Benzo[a]pyrene (BaP) is an environmentally relevant carcinogenic and endocrine disrupting compound that causes multigenerational effects in mammals. We hypothesized that, like in humans, BaP exposure would adversely affect zebrafish gene expression, reproduction and cause quantifiable pathologies across generations. Adult zebrafish (2 females x 2 males, N=10 replicate tanks per treatment) were fed 2% body weight/day flake food treated with 0, 11.6, 110, 1086 μg BaP/g flake (equivalent to 0, 0.23, 2.2, and 22 μg BaP/g fish/day) for 22 days. Parental gonad pathology and reproductive success, and F1 and F2 survival and morphological ab normalities were measured. The total number of eggs produced and fertilization success was non-significantly reduced in a dose-dependent manner, and parental ovarian atresia was significantly decreased. Mortality was significantly increased in F1 larvae whose parents were exposed to 2.2 and 22 μg BaP/g fish by 48 hours post-fertilization (hpf), respectively. High dose BaP F1 fish hatched sooner (48 hpf) compared to control (56 hpf). Body and tail shape and swim bladder were negatively impacted after parental exposure to 2.2 and 22 μg BaP/g fish. Moreover, differential gene expression by RNA-Seq of embryos and larvae was documented after a seven day parental embryonic water-borne BaP exposure. As expected, CYP1 genes in 96 hpf BaP-exposed embryos were induced as was vfg7. Based on these results, BaP negatively impacted zebrafish reproduction and development. Supported by NIEHS R01ES019940.

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Molecular Pathway of Neurotoxicity Caused by Developmental Exposure to Bisphenol A in Embryonic Zebrafish.

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Bisphenol A (BPA) is an endocrine disrupting compound widely used in consumer product manufacturing. Developmental exposure to 0.1 μM BPA results in a hyperactive phenotype in zebrafish larvae and impairment of learning in adults. The mode of action underlying these effects presumably involves impacts on nervous system development through activation of classical estrogen receptor (ERs) or other receptors such as estrogen related receptor gamma (ERR3) and G-protein coupled estrogen receptor (GPER). Transient knockdown of ERR3 using an anti-sense morpholino rescued the hyperactive phenotype in the zebrafish larvae suggesting a role of ERR3 in the developmental toxicity induced by BPA. We further used global miRNA and miRNA expression analysis to investigate transcriptional effects of BPA exposure and to identify candidate genes and signaling pathways that mediate the observed behavioral response. Zebrafish embryos were exposed to either of 0.1% DMSO, 0.1 μM BPA, 0.1 μM 17β-estradiol, 0.1 μM GSK4716 (an ERR3 agonist) from 8 hours post fertilization (hpf) to 24 hpf. At 24 hpf RNA was isolated to identify transcriptional changes that precede behavioral responses using a 135K NimbleGen microarray platform. Functional analysis of differentially expressed genes revealed CREB and prothrombin activation as top canonical pathways impacted by both BPA and E2 exposure, and suppressed expression of key genes involved in nervous system development and function. For miRNA expression analysis, total RNAs isolated at 24 hpf after exposure to either 0.1% DMSO or 0.1 μM BPA were labeled and hybridized onto an Esignature miRCURY LNA miRNA array (v. 11). The expression of a number of miRNAs was altered by BPA exposure. These results provide insight into potential modes of action underlying BPA’s neurodevelopmental toxicity in zebrafish. Supported by NIEHS T32 ES07060, R21 ES018970, P30 ES00210, P42 ES016465.

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Methylmercury-Induced Notch Signaling Points to a Role for Muscle Targets in Motor Nerve Development in Drosophila.

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Methylmercury (MeHg) is a ubiquitous environmental toxicant that targets the developing nervous system. Previous work in mammalian and insect cells has shown that MeHg induces expression of Notch pathway target genes, notably the Enhancer of split m-delta (E{spl}m{delta}) gene. We have shown earlier that MeHg induction of E{spl}m{delta} can occur independent of the Notch receptor and thus may directly influence neuronal development. In this study we examine the effects on MeHg on E{spl}m{delta} gene expression in parallel with neural development events in the Drosophila embryo. We now show that E{spl}m{delta} is specifically upregulated in MeHg-exposed embryos. We exclude the possibility that MeHg-induced E{spl}m{delta} expression is a by-product of a general stress response or a shift in developmental timing. MeHg phenotypes are apparent in the outgrowth of the embryonic intersegmental and segmental motor nerves. Genetic manipulations causing overactivity of the Notch pathway in neurons can mimic these phenotypes. Unexpectedly, induced expression of E{spl}m{delta} in neurons does not cause a failure of motor nerve outgrowth. We now demonstrate that endogenous E{spl}m{delta} expression localizes to developing muscle and that E{spl}m{delta} overexpression in embryonic muscle causes a segmental nerve phenotype similar to MeHg treatment. Closer examination shows altered patterning in muscle fields stemming from either MeHg treatment or E{spl}m{delta} overexpression. In contrast, targeting expression of the closely related E{spo}m{delta} to developing muscle shows no embryonic phenotype, whereas E{spo}m{delta} targeted to neurons is embryonic lethal. In summary these data highlight a novel mechanism whereby MeHg can engage the activity of a Notch pathway target gene to alter coordinated development of muscles and motor neurons.

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Effects of Benzo[a]pyrene on Early Zebrafish Development.


Benzo[a]pyrene (BaP) is a ubiquitous environmental contaminant that is an endocrine disrupting and carcinogenic high molecular weight polycyclic aromatic hydrocarbon. Our previous work found that BaP significantly decreased fish brain aromatase (CYP19b) expression, a key enzyme in steroidogenesis. We hypothesized that BaP deregulates the steroid hormone hypothalamus-pituitary-gonad feedback loop adversely affecting reproductive development and physiology. Zebrafish embryos were exposed to waterborne concentrations of BaP (0, 10, and 50 μg/L) for 96 hours postfertilization (hpf). Fifty μg/L BaP significantly increased mortality compared with the control and 10 μg/L groups at 24, 48, 72, and 96 hpf, whereas mortality was not significantly increased until 96 hpf in the 10 μg/L BaP group. In order to quantify effects on involved in xenobiotic and endobiotic metabolism, larvae were collected at 48, 72, 96, 168 and 504 hpf. Histopathological assessment of gonad maturation was done on paraffin embedded and sectioned fish at 28, 32, 35, and 52 days post fertilization. In a treatment-blinded morphological assessment of larvae at 96 hpf, the high BaP dose significantly decreased the body length, optic vesicle, and swim bladder size while increasing pericardial and abdominal edema.

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compared to control and 10 μL treatments. Body and tail shape and fin malformation scoring also indicate a dose-dependant adverse impact of BAP-exposure on development. Results extend previous studies highlighting the adverse impacts on early development of BAP-exposure. (Supported by NIEHS R03 ES018962)

332 Drosophila CYP6g1 and Its Human Homolog CYP3A4 Confer Tolerance to Methyl mercury during Development.
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The fetal nervous system is a well-known primary target for methyl mercury (MeHg) toxicity. Despite knowledge of numerous cellular processes that are affected by MeHg, the mechanisms that ultimately influence tolerance or susceptibility to MeHg in the developing fetus are not well understood. Using transcriptomic analyses of developing brains of MeHg tolerant and susceptible strains of Drosophila, we previously identified members of the cytochrome p450 (CYP) family of monoxygenases/oxidoreductases as candidate MeHg tolerance genes. CYP genes encode Phase I enzymes best known for xenobiotic metabolism in the liver as well as synthesis and degradation of essential endobiotics, such as hormones and fatty acids, that are critical to normal development. We now demonstrate that natural and induced variation in expression CYP genes can strongly influence MeHg tolerance in the developing fly. We show that modulating expression of a single CYP, CYP6g1, specifically in neurons or the fat body (liver equivalent) is sufficient to rescue Drosophila development in the presence of MeHg in the diet. Furthermore, we identify CYP3A4 as a human homolog of CYP6g1 and show that it similarly confers MeHg tolerance when ectopically expressed in flies. Finally, pharmacological induction of endogenous CYPs with caffeine also results in elevated tolerance to MeHg in developing flies. These findings establish a previously unidentified role for CYPs in modulating MeHg toxicity and point to a potentially conserved role of CYP genes to influence susceptibility to MeHg toxicity across species.

333 Systems Approaches to Define the Developmental Toxicity of Polybrominated Diphenyl Ethers.

Polybrominated diphenyl ethers (PBDEs) are high production volume flame retardants used in a number of consumer products. PBDEs are ubiquitous environmental pollutants due to their widespread usage, persistence and lipophilicity. Developmental exposure to PBDEs is associated with a number of neurological and developmental effects in humans and wildlife. A major complexity in assessing the hazard and risk posed by PBDEs is the diversity of chemical congeners; chemical structure-activity relationships are not established for these compounds. Our hypothesis was that developmental toxicity of PBDEs is highly structure dependent and that the molecular targets of these compounds are distinct. We performed rapid throughput assessment of PBDE developmental and neurotoxicity in zebrafish with PBDE congeners 47, 77, 99, 100, 153, 154, and 183. All exposures were from 6 until 120 hours post fertilization (hpf) at which time larvae were assessed for photo-induced locomotor activity and changes in a suite of 20 morphological endpoints. Preliminary global gene expression data suggested that some PBDE congeners may activate the aryl hydrocarbon receptor (AhR). Using the embryonic zebrafish model to rapidly assess developmental toxicity, global transcriptional responses and AhR activation in embryos exposed to parent, oxygenated PBDEs (OPAHs) and environmental mixture samples during development. Using comparative analysis of mRNA expression profiles from microarrays with embryos exposed to benzo(a)anthracene (BAA), dibenzo[b,k]pyrene (DB[b,k]P) and pyrene (PYR), we identified expression biomarkers and disrupted biological processes that precede developmental abnormalities. These transcriptional responses were associated with PAH body burdens in the embryos detected by GC-MS. We found that uptake data was essential for discerning molecular pathways from dose-related differences, and identified two primary toxicity profiles. While BAA disrupted transcript involved AhR signaling and vasculogenesis, DB[b,k]P and PYR misregulated ion homeostasis and muscle-related genes. Biomarkers of these toxicity pathways are under investigation with a diverse group of OPAHs, and comparative analysis of embryos exposed to OPAHs with different proposed molecular mechanisms using RNA-seq will expand and refine pathways of PAH-induced developmental toxicity. This research is supported by NIEHS grants P30ES00210, P42ES016465 and T32ES07060.

335 Defining Pathways of Polycyclic Aromatic Hydrocarbon Developmental Toxicity in Zebrafish Using Systems-Based Transcriptional Profiling.
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Polycyclic aromatic hydrocarbons (PAHs) are ubiquitous in the environment as components of fossil fuels and by-products of combustion. Defining toxicity mechanisms for this large family of multi-ring structures and substituted derivatives is a substantial challenge. In addition to the well-studied carcinogenic properties of several PAHs such as benzo(a)pyrene, reports of cardiac and developmental effects have increased concern about health risks of exposure to PAHs. Some PAHs induce toxicity via activation of the aryl hydrocarbon receptor (AhR), while others act through uncharacterized AhR-independent pathways. We employed the zebrafish model to rapidly assess developmental toxicity, global transcriptional responses and AhR activation in embryos exposed to parent, oxygenated PAHs (OPAHs) and environmental mixture samples during development. Using comparative analysis of mRNA expression profiles from microarrays with embryos exposed to benzo(a)anthracene (BAA), dibenzo[b,k]pyrene (DB[b,k]P) and pyrene (PYR), we identified expression biomarkers and disrupted biological processes that precede developmental abnormalities. These transcriptional responses were associated with PAH body burdens in the embryos detected by GC-MS. We found that uptake data was essential for discerning molecular pathways from dose-related differences, and identified two primary toxicity profiles. While BAA disrupted transcript involved AhR signaling and vasculogenesis, DB[b,k]P and PYR misregulated ion homeostasis and muscle-related genes. Biomarkers of these toxicity pathways are under investigation with a diverse group of OPAHs, and comparative analysis of embryos exposed to OPAHs with different proposed molecular mechanisms using RNA-seq will expand and refine pathways of PAH-induced developmental toxicity. This research is supported by NIEHS grants P30ES00210, P42ES016465 and T32ES07060.

336 Ebselen As a Countermeasure for Nitrogen Mustard Vescicant-Induced Toxicity.
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Mechlorethamine (HN2), a nitrogen mustard vesicant, is a bifunctional alkylating agent commonly used as a model to study sulfur mustard-induced lung injury. Previously, we reported that HN2 selectively targeted antioxidants in lung A549 epithelial cells including the selenoprotein thioredoxin reductase (TrxR) forming intra- and inter-molecular cross links. This resulted in dimer and oligomer formation, and enzyme inactivation, a process contributing to oxidative stress and toxicity. In this study, we examined the effects of ebselen, a selenium-containing antioxidant, on HN2-induced toxicity. In A549 cells, HN2 was found to be cytotoxic...
(LD50 = 25 μM). Both pre- and post-treatment with ebselen (50 μM) significantly attenuated HN2-induced toxicity. This was correlated with reduced inhibition of TrxR enzyme activity and decreased formation of TrxR dimers and oligomers. Ebselen treatment was found to induce TrxR protein expression in A549 cells, suggesting that inhibition of toxicity was due to both reduced TrxR damage and increased TrxR enzyme activity. Using purified rat liver TrxR, ebselen also protected against enzyme inactivation by HN2. This is due to a decrease in HN2 binding to the catalytic residues on TrxR, thus preventing enzyme inactivation. Taken together, our data suggest that ebselen has the potential to be an effective countermeasure for HN2-induced lung injury. Support: NIH grants AR055073, ES004738, CA132624, ES05022 and GM034310.

337 Identification of Thioredoxin As a Molecular Target for Sulfur Mustard Analog Vescants.

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The thioredoxin system, composed of thioredoxin reductase, thioredoxin and NAPDH, is a cellular disulphide reduction system important in antioxidant defense and cell growth control. We reported previously that thioredoxin reductase is a target of both monofunctional [1,2-chloroethyl ethyl sulfide (CEES)] and bifunctional (HN2, nitrogen mustard) voicing agents. These vesicants were found to covalently bind to selenocysteine/cysteine residues in the redox centers of the enzyme, leading to its inactivation and toxicity. Thioredoxin contains two catalytic cysteine and three structural cysteine residues. In these studies, we determined if vesicants also target thioredoxin. Both CEES and HN2 treatment were found to cause time- and concentration-dependent inhibition of thioredoxin in A549 cells. Western blot analysis revealed that a band corresponding to tetramers of thioredoxin was present in HN2-, but not CEES-treated cells, suggesting that HN2 caused inter-molecular thioredoxin cross-links. Using recombinant enzyme and mass spectrometry, we found that both CEES and HN2 alkylated cysteine and lysine residues on thioredoxin, including cysteine-32 and cysteine-35 in the redox centers of the enzyme. In addition, several inter-peptide cross links were identified in HN2-treated thioredoxin, providing a mechanism for the formation of thioredoxin tetramers in A549 cells. These data demonstrate that thioredoxin is a molecular target of sulfur mustard vesicants. Agents that can reactivate the thioredoxin system may be effective countermeasures for sulfur mustard-induced tissue injury. Support: NIH grants AR055073, ES004738, CA132624, ES05022 and GM034310.

338 Novel Therapeutic Compounds YEL001 and YEL002 Mitigate Radiation-Induced Toxicity Authors.

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The possibility of a radiation disaster from a nuclear detonation or accident has existed for over 50 years and spawned much of the basic research in radiobiology in the 1950-60s. The recent Fukushima accident was yet another reminder that there remains a dire need to develop novel therapies against radiation-induced toxicities. Here we report on the development of two novel radiation countermeasure therapeutics: Yel001 and Yel002. These small, biologically active, drug-like molecules were uncovered in the DEL high throughput assay reducing radiation-induced cytotoxicity and genotoxicity in yeast. Radiation-modulating activity was further confirmed in yeast plate-based DEL Assay; addition of either Yel001 or Yel002 to irradiated cultures reduced cell death and genomic instability. Further, Yel compounds increases survival to 75% in vivo following an LD100/30 dose of ionizing radiation (IR) with the first therapeutic injection administered 24 hours post exposure followed by injections at 48, 72, and 120 hours. Additionally, treatment with Yel001 and Yel002 compounds reduces radiation-induced leukemia from 90% to 20% and 40% respectively. Of note, treatment with either Yel001 or Yel002 reduced spontaneous leukemia rate from 10% to 0%. Treatment with Yel002 following IR accelerates the recovery of the hematopoietic cells after sub-lethal exposures. In addition, treatment with Yel002 reduces EMS, MMS, UV, cigarette smoke extract as well as nitrogen mustard induced toxicity as well as genotoxicity showing a broad application spectrum. Toxicity has not been observed in either in vitro or in vivo administrations. Overall, Yel compounds have much potential as stockpile therapies for radiation-induced lethality and cancer; they are highly effective when administered up to 24 hours post exposure, they reduce radiation-induced sequelae such as leukemia, and appear to have an acceptable toxicity profile.

339 Show Bispyridinium Nonoximes Direct Interactions with Muscle-Type Nicotinic Acetylcholine Receptors?

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Objective: In poisoning with some organophosphorus nerve agents, e.g. soman, therapeutic efficacy of oximes is limited. For such cases, a direct intervention at nicotinic acetylcholine receptors (nAChRs) might be an alternative. Studies with the bispyridinium non-oxime MB327 (1.1’-(propene-1,3-diyl)bis(4-tert-butylylpyridinium) diiodide) demonstrated a therapeutic effect against soman in vitro and in vivo. As MB327 was found to interact most probably with muscle-type nAChRs, improved therapeutic efficacy could possibly be achieved with compounds that show enhanced activity. To identify potential candidates, homologous series of substituted and non substituted analogues (linker C1-C10) of MB327 were investigated in binding and functional assays. In addition, their inhibitory activity was assessed with human acetylcholinesterase (AChE).

Experimental procedures: In competition radioligand binding assays, the influence on [3H]epibatidine binding sites of Torpedo californica nAChR was investigated. Functional assessments were performed with cell-free electrophysiology based on solid supported membranes (SSM), AChE inhibitory properties of the compounds were assayed with a modified Ellman assay using haemoglobin-free human erythrocyte ghosts.

Results: MB327 and several bispyridinium structure analogues exhibit no regular displacement curves at [3H]epibatidine binding sites. Compounds with unsubstituted pyridinium ring and long linkers (> C7) show regular competition but no intrinsic effect. Inhibition of human AChE (IC50 < 1 μM) for both bispyridinium compound series (unsubstituted and with p-tert-butylyl in both pyridyl rings) was observed with increasing distance between the pyridinium N (> C9 and > C6 respectively).

Conclusion: The interaction with [3H]epibatidine binding sites and functional improvement of Torpedo californica nAChR was dependent on the substitution and C-linker between the pyridine ring. Further research is necessary for better understanding how these compounds interact with the nAChR.

340 Novel Pyridinium Oximes Offer Neural Protection in the Central Nervous System against Nerve Agent Surrogates.

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Organophosphates (OPs), including nerve agents, target the cholinergic system via inhibition of acetylcholinesterases (AChE), with subsequent overstimulation resulting in neural damage and potential detrimental long-term effects. The efficacy of novel pyridinium oxime reactivators with moieties to increase blood-brain barrier penetration, was tested using highly relevant sarin and VX surrogates. Glial fibrillary acidic protein (GFAP; an indicator of neural damage) and monoamines (dopamine, serotonin (5-HT) and their metabolites) were measured in select brain regions via immunohistochemistry and HPLC, respectively. Adult male rats were treated ip with high, sub-lethal doses of surrogates for sarin or VX, nitrophenyl isopropyl methylphosphonate (NIMP; 0.325mg/kg) or nitrophenyl ethyl methylphosphonate (NEMP; 0.4mg/kg), respectively. Surrogate treatment was followed after 1 hr by im administration of novel oxime (0.1mmol/kg). Seizure activity was monitored, and kainic acid (KA; 10mg/kg) served as a positive control. Administration of KA or surrogate (NIMP or NEMP) significantly (p<0.05) increased GFAP expression compared to control animals. Two different formulations of oxime (bromide vs. mesylate salt) attenuated seizures and reduced GFAP levels over NIMP or NEMP treatments alone to levels near those of controls (p<0.05) in both the piriform cortex and dentate gyrus region of the hippocampus, while 2-PAM did not provide protection. Serotonergic activity was also altered in several brain regions, including the piriform cortex (significant increase in both 5-HT and 5-HIAA; p<0.05), one hr after NIMP treatment, with early markers of oxidative stress (isoprostanes) also being tested. These results indicate the potential therapeutic efficacy of these oximes and suggest this innovative chemistry may protect against neural damage induced by OPs. Supported by Defense Threat Reduction Agency: 1.E0056_08_AHB_C.
Sulforaphane Induces Antioxidants and Glutathione S-Transferase A1 and Protects against Vesicant-Induced Toxicity in Mouse Keratinocytes.

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Sulfur mustard and the related skin vesicant nitrogen mustard (methylchloroethyl hydrochloride, HN2) are bifunctional alkylating agents known to cause oxidative stress and persistent tissue damage including blistering. In the present studies we determined whether HN2-induced oxidative stress and cytotoxicity in mouse keratinocytes could be mitigated by upregulating antioxidant enzymes. Sulforaphane (SFN), an isothiocyanate found in cruciferous vegetables including broccoli, is a well characterized inducer of antioxidants and phase II metabolizing enzymes through a pathway mediated by the transcription factor nuclear factor-erythroid 2-related factor 2 (Nrf2). Treatment of PAM212 mouse keratinocytes with HN2 inhibited keratinocyte growth (IC50 = 0.1 μM). Pretreatment of the cells with 3 μM SFN for 3 hours protected against growth inhibition (IC50 = 6.0 μM). SFN also protected primary mouse epidermal keratinocytes from HN2 (IC50 = 2.0 μM and 30 μM with and without SFN, respectively). SFN also decreased HN2-induced phosphorylation of histone H2AX, a marker of DNA damage, in PAM212 cells. SFN was functionally active causing Nrf2 to translocate from the cytosol to the nucleus. This was associated with a marked increase in expression of the antioxidants heme oxygenase-1, thioredoxin reductase 1, and NADPH quinone oxidoreductase-1, as well as the phase II metabolizing enzyme glutathione-S-transferase A1. Taken together, these data indicate that oxidative stress is important in HN2-induced toxicity in mouse keratinocytes and that SFN may be an effective countermeasure.

Inhibition of AChE following exposure to toxic OPs (nerve agents and pesticides), results in the overstimulation of the nervous system. The approved therapies include an oxime, e.g. 2-PAM, to reactivate OP inhibited AChE; however these oximes have little, if any ability to cross the BBB. A series of novel pyridinium oximes has been synthesized that incorporate moieties that increase BBB penetration and AChE reactivation. Oximes were screened in vitro for their ability to reactivate AChE inhibited by a sarin surrogate, phthalimidyl isopropyl methylphosphonate (PIMP), or a VX surrogate, nitrophenyl ethyl methylphosphonate (NEMP), which phosphorylate AChE with the same moiety as sarin or VX, as well as with paraoxon, the active metabolite of the insecticide parathion. Rat brain homogenate was incubated with a concentration of OP that yielded about 80% AChE inhibition, followed by an oxime (0.1 mM) and AChE activity measured. In vitro AChE reactivation varied among oximes but was similar for each of the nerve agent surrogates; PIMP, 14-79%, and NEMP 23-76%, but differed for paraoxon 16-93%. Oxime lipophilicities ranged from 0.009 to 2.244 (Kow) and were greater than for 2-PAM (0.006). A subset of oximes that demonstrated AChE reactivation ≥ 40% in vitro were selected for testing in vivo in rats with the nerve agent surrogates. A high sublethal dose of a stable sarin surrogate, nitrophenyl isopropyl methylphosphonate (NIMP) (0.325 mg/kg) or NEMP (0.4 mg/kg) was administered ip, yielding about 80% brain AChE inhibition, followed by an im injection (0.1 mmol/kg) of an oxime at the time of peak brain AChE inhibition (1 hr). Twelve of 25 novel oximes tested yielded 10-35% brain AChE reactivation and attenuated OP induced seizures, indicating their ability to cross the BBB, reactivate brain AChE and thus demonstrating their therapeutic potential. Supported by DTRA-1.E0056.08.AHB.C.
Diazepam is routinely used to control organophosphate-induced seizures. The present study was designed to evaluate the effectiveness of diazepam to control seizures induced by soman (GD), disopropylfluorophosphate (DFP) and paraoxon. Prior to experimentation, animals were instrumented with cortical electrodes for monitoring brain activity. Animals exposed to GD (180 μg/kg, sc) were pretreated with HI-6 (125 mg/kg, ip) 30 min prior to GD challenge and treated with atropine methyl nitrate (2.0 mg/kg, im). Animals exposed to DFP (4.0 mg/kg, sc) were pretreated with pyridostigmine (0.026 mg/kg, ip) 30 min prior to DFP challenge and treated with atropine sulfate (AS; 0.2 mg/kg, im) and 2-PAM (25 mg/kg, im). Animals exposed to paraoxon (1.05 mg/kg, sc) were treated with AS (2.0 mg/kg, im) and 2-PAM (25 mg/kg, im). In all cases, diazepam (10 mg/kg, im) was given at 40 min after the onset of seizures. Diazepam was effective in terminating DFP- and paraoxon-induced seizures, but it was ineffective in terminating GD-induced seizures. In addition, neuropathology in animals treated with diazepam at 40 min after DFP or paraoxon-induced seizures was markedly reduced at 24 hr after seizure onset. In contrast, diazepam treatment at 40 min after the onset of seizures induced by GD did not prevent or attenuate brain injury. DFP and paraoxon models were developed as surrogate animal models for nerve agent (NA) biomedical research for evaluation of putative therapeutics. The difference in the effectiveness of diazepam in controlling seizures and reducing neuropathology induced by GD, DFP or paraoxon indicates that DFP and paraoxon are not suitable simulants for NA biomedical research, at least not for screening of potential anticonvulsants to control on-going seizures. Novel anticonvulsants that are effective in terminating seizures induced by DFP or paraoxon may not be efficacious against NA-induced seizures.

### 347 Comparative Anticonvulsant Effects of Diazepam in Rats


Diazepam is routinely used to control organophosphate-induced seizures. The present study was designed to evaluate the effectiveness of diazepam to control seizures induced by soman (GD), disopropylfluorophosphate (DFP) and paraoxon. Prior to experimentation, animals were instrumented with cortical electrodes for monitoring brain activity. Animals exposed to GD (180 μg/kg, sc) were pretreated with HI-6 (125 mg/kg, ip) 30 min prior to GD challenge and treated with atropine methyl nitrate (2.0 mg/kg, im). Animals exposed to DFP (4.0 mg/kg, sc) were pretreated with pyridostigmine (0.026 mg/kg, ip) 30 min prior to DFP challenge and treated with atropine sulfate (AS; 0.2 mg/kg, im) and 2-PAM (25 mg/kg, im). Animals exposed to paraoxon (1.05 mg/kg, sc) were treated with AS (2.0 mg/kg, im) and 2-PAM (25 mg/kg, im). In all cases, diazepam (10 mg/kg, im) was given at 40 min after the onset of seizures. Diazepam was effective in terminating DFP- and paraoxon-induced seizures, but it was ineffective in terminating GD-induced seizures. In addition, neuropathology in animals treated with diazepam at 40 min after DFP or paraoxon-induced seizures was markedly reduced at 24 hr after seizure onset. In contrast, diazepam treatment at 40 min after the onset of seizures induced by GD did not prevent or attenuate brain injury. DFP and paraoxon models were developed as surrogate animal models for nerve agent (NA) biomedical research for evaluation of putative therapeutics. The difference in the effectiveness of diazepam in controlling seizures and reducing neuropathology induced by GD, DFP or paraoxon indicates that DFP and paraoxon are not suitable simulants for NA biomedical research, at least not for screening of potential anticonvulsants to control on-going seizures. Novel anticonvulsants that are effective in terminating seizures induced by DFP or paraoxon may not be efficacious against NA-induced seizures.

### 348 Determination of Cholinesterase Levels in the Blood and Brain following Galantamine Administration in the Guinea Pig Model


The CounterACT Program at the National Institutes of Health is investigating galantamine hydrobromide (GAL) as a treatment for organophosphorous nerve agent (NA) intoxication and is establishing a dose response representation to determine optimal doses for future efficacy studies. GAL is a reversible acetylcholinesterase (AChE) inhibitor that crosses the blood–brain barrier and is presently approved for the treatment of mild to moderate Alzheimer’s disease. It has shown some efficacy against NA toxicity in animal studies. Ongoing studies are investigating the efficacy of GAL when administered either as a pre-treatment or post-treatment and when used in conjunction with standard NA treatments atropine and 2-PAM (pralidoxime). The objective of this study was to correlate clinical biomarkers and pharmacodynamics with clinical observations after treatment with GAL. Blood and brain levels of GAL and ChE inhibition were determined at various time-points along with standardized clinical observations. Cohort A used 36 animals with six dose levels of 0, 2, 4, 8, 12 and 16 mg/kg and serial blood collections from 5 minutes to 24 hours post treatment. These samples were assayed to determine the percent AChE activity at seven time-points. Cohort B used 144 male Hartley guinea pigs with six dose levels of 0, 2, 4, 8, 12 and 16 mg/kg with designated sacrifice time-points on 4 animals for each dose level at 5, 30, 60 min, 2, 4 and 24 hours post treatment. Clinical observations showed a trend of increasing severity of clinical observations as the dose level increased. All clinical signs started at 5 minutes post treatment and 2 animals were normal by the 24 hour post treatment time-point. Significant observations of toxicity were present at and above the 12 mg/kg dose level, combined with the greatest inhibition of AChE. Inhibition of AChE was seen at the 5 minute time-point (post treatment) with results showing a dose dependent inhibition from low to high dose levels.

### 349 Characterization of Protein Adducts of Nitrogen Mustards As Potential Exposure Biomarkers

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Protein adducts are useful as longer-term biomarkers of exposure to electrophilic xenobiotics, including chemical warfare agents (CWA) and their metabolites. The purpose of this study was to characterize protein adducts of the nitrogen mustards

### 345 Prediction of Human Equivalent Dose (HED) of Levofloxacin (LEVO) in New Zealand White Rabbits (NZW) to Support Efficacy of Combination Treatment with Novel Therapeutics against Inhalation Anthrax (IA).

M. Beliveau1, A. L. Menard1, J. F. Marier1, J. T. Troyer2 and E. K. Leftel2.

Regulatory agencies have suggested that novel therapeutics for treatment of IA do not diminish the efficacy of the standard-of-care (SOC) antibiotics in combination therapy, which would be the likely treatment. The NZW is an accepted animal model for IA where the SOC LEVO is both well tolerated, efficacious and where benefits of combination therapy would therefore need to be demonstrated. However, for proper demonstration of the added benefit of the new drug, the administered dose of LEVO must not be supra-therapeutic. Therefore, a HED of LEVO in NZW must be used. The objective of the current analysis was to determine pharmacokinetic (PK) parameters of LEVO in NZW and to determine the NZW dose equivalent to the HED of 7.14 mg/kg (500 mg QD) for IA. LEVO was administered to NZW by either PO or IV bolus administration and blood for PK analysis was collected at various times up to 48h post dose. Plasma samples were assayed for concentrations of LEVO using a validated LC/MS/MS method. Following calculation of PK parameters such as clearance (CL) and volume (Vss) by standard non-compartmental methods, the NZW HED was extrapolated from the clinical dose using the following equation: Dose_{rabbit} (HED) = (HED/CL_{human}) x CL_{rabbit} where the HED = 500 mg and the CL_{human} was taken from the literature (175 mL/min). Following IV dosing, mean CL_{rabbit} and Vss of LEVO were 17.7 mL/min/kg and 1619 mL/kg, respectively. Bioavailability in NZW was dose dependent and ranged from 37.9% to 79.7%. Using CL_{rabbit}, CL_{human} and the HED, the dose of LEVO in NZW corresponding to the HED was 50.5 mg/kg. If administered PO, the NZW HED would range from 53.8 mg/kg to 63.1 mg/kg. In conclusion, the above results suggest that LEVO IV dosing of 50.5 mg/kg or PO dosing of 53.8 to 63.1 mg/kg in NZW in combination with novel anthrax therapeutics are expected to allow sufficient demonstration of any added benefit of the novel drug.

### 346 The Beneficial Effect of a Treatment with Macrophages on Sulfur Mustard Cutaneous Burns.

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Sulfur mustard (HM) is a potent vesicant that its toxicology has been for many years a subject of research, yet, the exact mechanism of its toxicity is still elusive and treatment is only partially effective. Macrophages are known to play an essential role in almost every stage of wound healing and there is evidence for their beneficial effects in treating decubital ulcers and deep sterna wound infections in human. This study was aimed to investigate the efficacy of a treatment with activated macrophages in ameliorating acute and long-term HD induced skin injuries in the hairless guinea pig (HGP) model. HGP were exposed to HD vapor creating superficial dermal skin lesions. They were treated with either a single or multiple intra-dermal injections of human activated macrophages (hAMS) into the wound bed. Clinical and histological evaluations were conducted up to four weeks post-exposure. A single intra-dermal injection into the wound bed administered early (15min or 6hr) after exposure inflicted an initial positive effect on the extent of the damage, demonstrating a temporal decrease in clinical symptoms, reduced number of acidoophilic epithelial cells and less micro-vascations compared to untreated control. Repeated injections with hAMS (15min, 48hr and 7d post-exposure), decreased significantly the wounds’ area and improved the integrity of the barrier function as expressed by trans-epidermal water loss measurements up to 10 days. Still, at 21 days there were no differences between the experimental groups. The results demonstrated a beneficial effect of macrophages on HD induced lesions. Further investigation is required to determine whether macrophages are required during the early phase of wound development or during the late phase of scar formation and remodeling.
Military and civilians exposed to Gulf War-related stressors experienced higher rates of somatic symptoms than their non-exposed counterparts. The study was conducted at the University of Tbilisi, Tbilisi, Georgia.

We observed a significant increase in somatic symptoms among the exposed group compared to the non-exposed group. The results suggest that exposure to stressors during the Gulf War has long-term effects on the physical health of the exposed population. Further research is needed to understand the mechanisms underlying these effects and to develop effective interventions to mitigate their impact.
Soman (GD) is a potent chemical warfare nerve agent (CWNA). GD irreversibly inhibits acetylcholinesterase (AChE), resulting in accumulation of the neurotransmitter acetycholine (ACh). High ACh at synapses results in prolonged stimulation of muscarinic and nicotinic receptors leading to cholinergic crises. This study developed a novel head-out inhalation exposure system to serve as a realistic and applicable model for a mass casualty-type scenario to study the development of acute respiratory and neurological toxicity. GD vapor (520, 560, 600 and 825 mg/m^3) was generated in a saturator cell and carried by filtered N2 into a customized glass exposure chamber. Male rats (250-300 g) were restrained in head-out exposure tubes and exposed to GD for 4-10 minutes. The prebit analyzed LC50 of vaporized GD was 593.1 mg/m^3. A majority of the animals developed severe cholinergic responses followed by convulsions and died within 4-8 min post-exposure (PE). Mild to severe convulsions and muscular fasciculations were observed up to 24 h in animals exposed to 560 and 600 mg/m^3. GD exposures produced significant, concentration-dependent inhibition in cardiac blood AChE activity. AChE activity was inhibited in bronchoalveolar lavage (BAL) fluid, lung and whole brain tissues. Rats exposed to a 600 mg/m^3 exhibited severe brain damage in the piriform cortex, neocortex, dentate hilus, amygda and thalamus. Analyses of lung tissue showed morphologic changes in alveolar histiocytes, hemorrhage and inflammation consisting of neutrophilic exudate. Respiratory and neurological toxicity induced by GD vapor was determined by clinical observations, histopathology and biochemical analyses of tissues 24 h PE. We integrated novel technologies in inhalation toxicology and physiology to evaluate CWNA toxicity to develop an improved model suitable for testing medical countermeasures in a mass casualty scenario. This research was supported by the Defense Threat Reduction Agency project #3F0014.
miclal forms of TrxR and Trx. HN2 was significantly more efficient in inhibiting cytotoxic, when compared to mitochondrial forms of the enzymes. For TrxR1 and TrxR2, the IC50’s for inhibition of enzyme activity were 2.7 μM and 14.3 μM, respectively. Immunoblot analysis of subcellular fractions from control cells on denaturing SDS-PAGE showed TrxR (57 kDa) and Trx (14 kDa) monomers. HN2 treatment cross-linked Trx1 and Trx2 forming tetramers, and TrxR1 and TrxR2 forming dimers and oligomers. These data demonstrate that HN2 can target the thioredoxin system in cytosolic and mitochondrial fractions of A549 cells by cross-linking proteins. Disruption of this system in lung cells provides a mechanism for nitrogen mustard-induced oxidative stress and toxicity.

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360 Dose Modulation of a Potential Therapeutic for Neovascularization from Ocular Exposure to Sulfur Mustard.

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Sulfur mustard is a potent alkylating agent affecting the skin, respiratory tract and eyes. Ocular exposure to HD can result in long term injuries including corneal neovascularization (NV) and blindness. The immunosuppressant, cyclosporine A, has shown promise for protecting epithelial cells against apoptosis in in vitro experiments. To investigate the effects of cyclosporine A in vivo, two studies were conducted, an initial long term study for 112 days, and a 28 day study using the white rabbit ocular model to evaluate the effects of different concentrations of cyclosporine A. The right cornea was insulted with 0.4 μg/kg of HFA, which turned to baseline by 28 day. In contrast H3K4 methylation decreased 1 d - 7 d post-NM; this was followed by an increase at 28 d. Additionally, whereas H3K36 methylation increased beginning 3 d post-NM and persisted up to 28 d, increases in H3K27 methylation were only evident 28 d after NM exposure. These data suggest that NM-induced lung injury leads to specific alterations in chromatin structure which may contribute to pro-inflammatory and antioxidant gene expression. Supported by NIH Grants AR055073, ES004738, CA132624, GM034310, and ES005022.

361 Development and Optimization of In Vitro Models of Chemical Ocular Injury for High-Throughput siRNA Screening.

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Chemically induced ocular injuries are considered one of the few true ocular emergencies based on their high potential to inflict rapid and significant tissue damage. Currently there are no specific therapeutics to treat chemically induced corneal injury, and there is a pressing need to rapidly screen therapeutics to discover a treatment. Understanding of the molecular mechanisms of this injury is necessary to aid in rational therapeutic development. We intend to utilize high throughput small inhibitory RNA (siRNA) screening to elucidate the mechanisms of chemical cornea injury and to identify therapeutic targets. Herein we present the development of the in vitro models for this screening effort. Two immortalized human corneal epithelial cell lines will be used for screening: SV40 large T antigen immortalized corneal epithelial cells (SHECs) and telomerase immortalized corneal epithelial cells (TCECs). Hydrofluoric acid (HFA) is used for the induction of chemical injury. The major conditions optimized included cell line plating density, exposure dose, and transfection conditions. Matrices of these conditions were evaluated to ensure optimal target knockdown without sacrificing cell viability. Transfection conditions were optimized using siRNA targeting cyclophilin b, and knockdown was assessed using the QuantiGene Plex assay, a bead based assay using hybridization to amplify specific miRNAs allowing for quantification. A dose-response study of HFA injury was performed, and the HFA-induced production of the cytokines was evaluated by multiplex bead-based assay for a panel of 30 cytokines. Once optimized, these models will be used to screen subgenomic siRNA libraries targeting HF-induced ocular inflammation.

Disclaimer: The views expressed in this article are those of the author(s) and do not reflect the official policy of the Department of Army, Department of Defense, or the U.S. Government. This research was supported by the Defense Threat Reduction Agency – Joint Science and Technology Office, Medical S&T Division.

362 Multiarray Gene Expression Approach Analyzing Cultured Primary Human Astrocytes and Neurons following Chemical Warfare Nerve Agent Exposure.

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Chemical warfare nerve agents (CWNAs) are irreversible organophosphorus cholinesterase inhibitors that cause seizure. Within 24 h of CWNAs exposure, neuronal degeneration, astrocytic activation and cellular death can be observed in various brain regions. In this study, we investigated the effects of CWA exposure on primary cultures of neurons and astrocytes to determine direct molecular alterations in the absence of CWA-induced seizure activity. Cultured primary human astrocytes were exposed to 100 μM of the CWNAs sarin, soman, VX, and harvested at 24 h. Similarly, cultured primary human neurons were exposed to 100 μM soman or VX and harvested at 24 h. Total RNA was isolated from exposed cells, processed for microarray analysis, and hybridized to Affymetrix Human Genome U133 Plus 2.0 arrays. Principal component analysis (PCA) of the astrocyte data did not show distinct profile differences based on agent type, indicating few if any significant differences in transcript expression in response to agent exposure. However, PCA of the neuronal data showed a distinct profile difference based on agent type. For both cell types, the dataset was filtered by agent type, and an analysis of variance (ANOVA) was performed using agent as the factor. Using these combined data, the pathways significantly affected by agent type in cultured human primary neurons were identified. The genes differentially expressed in response to soman exposure mapped to canonical pathways containing inflammatory molecules. Using the same samples, real-time PCR was utilized to provide quantitative measurements of gene transcription across genes associated with the human immune response. These data suggest a non-cholinergic effect of soman that induces transcriptional expression of inflammatory molecules as well as components of the GABAA receptor pathway. In addition, these data suggest that an in-depth analysis of in vivo transcriptomic data focusing on the GABA pathway may reveal new targets within that pathway for improved medical countermeasures to these toxic agents.
MicroRNA Microarray Analysis of Human-Induced Pluripotent Stem Cell-Derived Neurons and Cardiomyocytes following Exposure to the Organophosphate Nerve Agents Soman and VX.


Chemical warfare nerve agents (CWNA) are potent cholinesterase inhibitors that may also have non-cholinesterase effects. Several in vitro studies have demonstrated CWNA-induced damage in the brain and heart following CWNA exposure. The mechanisms of this damage have been a critical area of research for the development of medical countermeasures. This study utilized microRNA (miRNA) analysis to evaluate potential direct cellular effects of the nerve agents soman (O-Pinacolyl methylphosphonofluoridate) or VX (o-ethyl-s-[2 (diisopropylamino) ethyl]) on human-induced pluripotent stem cell (iPSC)-derived neurons and cardiomyocytes. This approach was taken since miRNA expression changes are stimulus specific and no previous studies of miRNA profiles have been conducted for CWNA exposure. Cells were treated with soman or VX at concentrations of 0 μM, 0.1 μM or 100 μM for either 1 h or 6 h. Following treatment, isolated total RNA was processed for miRNA microarray analysis and analyzed for significant changes. Soman- and VX-treated samples were analyzed separately. Principal component analysis (PCA) was used to identify major sources of variability in the dataset. PCA analysis of neurons identified differences. miRNA expression only for RNA isolated at 6 h to soman. Targets that were significantly altered under these conditions were miR-2277, miR-1910 and miR-1972. miR-2277 was significantly altered in soman- and VX-exposed neuron data sets that were analyzed by both time and dose. Minimal sample variability was observed with cardiomyocytes as determined by PCA analysis. One-way ANOVA with time as the factor identified miR-3178 as the only target that was altered significantly by both soman and VX in cardiomyocytes. This miRNA modulates several targets including complexin-1 and splicing factor-1. This study demonstrates the feasibility of using miRNA microarray analysis for the study of CWNA cellular effects.

Toxidromes—A Decision-Making Tool for Early Response to Chemical Mass Exposure Incidents.

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A common language to describe and recognize clinical manifestations of toxic chemical exposures is essential for emergency responders and hospital first receivers to be prepared to provide rapid and appropriate medical care for victims of industrial chemical mass exposures and terrorist attacks. In these situations, when the identity of the chemical is not known, first responders need a tool to rapidly evaluate victims and identify the best course of treatment. Military and civilian emergency responders use a "toxic syndrome" (toxidrome) approach to quickly assess victims and determine the best immediate treatment when information on chemical exposures is limited. Toxidromes can be defined by a unique group of clinical observations, such as vital signs, mental status, pupil size, mucous membrane irritation, and lung and skin examinations. Data on over 20 toxidrome systems were evaluated to identify salient features and develop a consistent lexicon for use by state, local, tribal, territorial, and federal first responders and first receivers. A workshop of over 40 practitioners and experts in emergency response, emergency medicine, and medical toxicology developed names and definitions for 12 unique toxidromes that describe and differentiate the clinical signs and symptoms from exposures to chemicals. These toxidromes focus on acute signs and symptoms caused by inhalation and dermal exposures. Each toxidrome is characterized by exposure routes and sources, organs/systems affected, initial signs and symptoms, underlying mode of action, and treatment/antidotes. Toxidrome names and definitions are designed to be readily understood and remembered by users. Communication in a crisis requires accurate and succinct terms that can quickly convey the health conditions of patients. These toxidromes lay the foundation for a consistent lexicon, that if adopted widely, will improve response to chemical mass exposure incidents.

Progression of Injury and Toxic Effects following Nitrogen Mustard Exposure in SKH-1 Hairless and C57BL/6J Mice Skin: Clinical and Pathological Significance.


Lack of availability of relevant skin injury models and clinically applicable biomarkers are major limitations in developing therapies to rescue skin injury and vesicant by chemical warfare agent sulfur mustard (SM). Consequently, we conducted studies to establish useful clinical and histopathological endpoints with primary vesicating agent nitrogen mustard (NM). NM possesses strong vesicating and alkylating properties and causes damage to cellular macromolecules, exerting severe skin toxicity comparable to SM. Our comprehensive studies employing NM (3.2 mg) exposure for 12, 24, 72 and 120 h in both SKH-1 and C57BL/6J mice showed clinical sequelae of toxicity, including visible microblistering (12–24 h), edema (12–120 h), erythema (1.2–24 h), and hyper- and hypopigmentation, wound healing and scaly dry skin (72–120 h) that were comparable in both the mouse strains, and similar to those reported with SM in humans. In addition, 40% mortality was observed by 120 h after NM exposure in C57BL/6J mice. H&E stained skin sections of both mice showed that NM (12–120 h recovery) caused increased skin bi-fold thickness; histopathological effects such as hyperproliferation, microcresication, epidermal and dermal necrosis, denuding and scab formation, and parakeratosis (24–120 h), hypercornification (12–120 h), acanthosis and re-epithelialization (72–120 h); increase in inflammatory cells; and red blood cell extravasation into the dermis. These histopathological effects with NM were comparable to those reported in humans and other animal species with SM, and were quantified as percent of skin sections showing these effects, fold increases in histopathological abnormalities, and prevalence among mice (% that showed these effects). These NM-induced effects are novel clinically relevant biomarkers to be used in screening and optimization of rescue therapies for skin injuries due to NM and SM in humans.

Activation of DNA Damage Repair Pathways in Response to Nitrogen Mustard-Induced DNA Damage and Toxicity in Skin Keratinocytes.

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Alkylating agent nitrogen mustard (NM), a structural analog of chemical warfare agent sulfur mustard (SM), upon exposure to tissues, forms adducts and crosslinks with DNA, RNA and proteins. The major mechanism of NM-induced toxicity involves DNA interstrand crosslinks (ICL) formation resulting in either induction of cell cycle arrest to facilitate DNA damage repair or cell death, in the case of inade-quate repair. Consistently, NM (0.75 μM) exposure in mouse epithelial JB6 cells decreased cell growth and caused S-phase arrest by 16 h after exposure. Cells were then released to G2-M phase by 24 h and resumed normal cell cycle progression by 48 h after exposure. Repair of NM-induced DNA ICLs involves formation of DNA double strand breaks (DSB). Our studies showed an increase in comet tail extent moment starting between 4 and 8 h of NM exposure, as well as an increase in levels of DNA DSB repair molecules (phospho H2AX and p53, radi50 and XRCC1). The repair of DNA double strand breaks occurs via homologous recombination repair (HRR) or through the non homologous end joining pathway (NHEJ). The activation of the HRR pathway was evidenced by formation of Rad51 foci at 4 h after NM exposure, and activation of NHEJ pathway was indicated by increases in phospho and total DNA-PK levels. To confirm this, NHEJ and HRR pathways were inhibited by using DNA-PK inhibitor NU7026 and Rad51 siRNA, respectively. Inhibition of NHEJ did not result in a significant decrease in total cell number after 48 h of NM exposure and also did not affect the NM-induced S-phase arrest at 16 h. However, inhibition of the HRR pathway caused a 28% decrease in cell number, and a lack of NM-induced S-phase arrest, probably leading to an increase in the observed cell death. These studies indicate that HRR may be a key pathway involved in repair of NM-induced DNA DSBs. These findings may be useful in developing new therapeutic strategies against NM-induced skin toxicity.
Sulfur mustard (HD) causes severe chemical burns to the skin, eyes, and airways. HD was used as a chemical warfare agent (CWA) in the Iran/Iraq conflict, and more than half of surviving HD-exposed casualties suffer from permanent lung injuries. The mechanisms and timing of the development of these pathologies are poorly defined, and there is no effective antidote. Rats were intubated and ventilated for 10 min with nebulized HD or vehicle to achieve doses of 0, 0.5, 1.75, 2.25, and 3 mg/kg. Pulmonary function was analyzed by whole-body plethysmography. Rats were euthanized at various time-points (6 months post-exposure), blood chemistry was analyzed, broncho-alveolar lavage fluid was analyzed for cytokines/redox state, and lungs were subjected to pathologic analysis. The data show high correlation between blood gas perturbations, upper airway necrosis and pulmonary function in the 24 h to 48 h after the 3 mg/kg HD exposure. In longer-ranged animals, the 3 to 7 wk period was a significant challenge because 15% of the 3 mg/kg group and 10% of the 2.25 mg/kg group died suddenly or required withdrawal from the study during this time. Alveolar exudates, edema, oxidative stress, and inflammation peaked at 3 wks and correlated with changes in pulmonary function and respiratory distress. Therefore, the first day and the second to third wks post-exposure may be crucial windows in the progression of HD inhalation injury, and treatment at these times may be the best approach. Based on this, we have evaluated single and multiple infusions of mesenchymal stem cells as a treatment and have found success in reducing HD-induced edema, necrosis, inflammation, and death at 5 wks post-exposure. This study provides the first long-term examination of HD-induced lung injury and systemic effects, and demonstrates the feasibility of stem cell therapies for treatment of HD inhalation injury.

Sulfur mustard (SM) is a primary vesicating warfare agent that upon exposure causes severe skin injuries. Currently, we lack effective antidote against SM-induced skin injuries, in part due to lack of appropriate animal model(s) that can be used for mechanistic and efficacy studies in laboratory settings. Our earlier studies have established biomarkers related to inflammation and vesication in SKH-1 hairless mice using 2-chloroethyl ethyl sulfide (CEES), an SM analog. However, CEES is a monofunctional alkylating agent that is less toxic than SM. Therefore, to develop a more relevant skin injury model, we have now used sulfur mustard (NM); a primary vesicant and a bifunctional alkylating agent that induces toxic effects comparable to SM. We compared the effect of NM (3.2 mg/kg) exposure for 12, 24, 72 and 120 h in SKH-1 hairless and C57BL/6J mice. NM caused significant increases in skin microvesication, cell proliferation, apoptotic cell death, inflammatory cells (neutrophils, macrophages, and mast cells) and MPO activity in both mouse strains. However, in SKH-1 mice there was a more prominent increase in epidermal thickness, macrophages and mast cell infiltration, relative to that seen in C57BL/6J mice. NM also caused collagen degradation at early time points (12-24 h) with a decrease in collagen trichrome staining and an increase in dermal thickening, due, at least in part, to edema. However, at later time points (72 and 120 h), dense collagen staining with reduced dermal edema was observed, indicating a healing process. This study indicates that both mouse strains have comparable susceptibility to NM injury, as shown by inflammation and vesication. However, some inflammatory responses were more pronounced in NM-exposed skin in SKH-1 mice. These newly established biomarkers in a more accessible and relevant NM skin injury model should aid in identifying effective therapies for treatment of skin injuries due to NM and SM.
Mechanisms of Vesicating Agent Nitrogen Mustard-Induced Skin Injury in SKH-1 Hairless Mice.

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Lack of a comprehensive understanding of molecular mechanisms and signaling pathways involved in skin injuries due to chemical warfare agent sulfur mustard (SM) is a major limitation in developing mechanism-based therapies for rescue of skin injuries by vesicating agents. Our recent studies in SKH-1 hairless mice show that exposure to nitrogen mustard (NM: 3.2 mg) for 12, 24, 72, 120 or 168 h triggers an inflammatory response, vesication, apoptotic cell death, and either initiation of or 40% mortality by 168 h of exposure. Both SM and NM vesicant-induced tissue injury is mainly due to alkylating properties, with DNA damage resulting from direct toxicity and/or oxidative stress. In this study we extended our efforts to identify the mechanisms involved in NM-induced toxic response at 12-120 h after exposure. In skin tissue of SKH-1 mice, NM exposure caused p53 ser15 phosphorylation and increased p53 accumulation, both indicating DNA damage. Our results also showed that NM exposure, with recovery for 12-168 h, induced expression of inflammatory mediators COX-2 and iNOS in mouse skin, and 2-fold increase in expression of protease MMP-9, that may contribute to NM-induced vesication. NM exposure, with recovery for 12-168 h, also was associated with phosphorylation of mitogen-activated protein kinases (MAPKs: ERK1/2, JNK, and p38) and increased oxidative stress, as indicated by enhanced 4-HNE (lipid peroxidation) and DMPO protein adduct formation. Our results thus far indicate that NM induces activation of upstream signaling pathways including MAPKs and oxidative stress that could be responsible for NM damage, cell death, and expression of inflammatory and proteolytic mediators contributing to the inflammatory response and vesication. These molecular targets could be useful in developing therapeutic interventions against NM- and SM-related skin injuries.

Ovarian Hormones Affect the Physiological Response to VX in Female Rats.


Chemical warfare nerve agents (CWNAs), such as VX, irreversibly bind to acetylcholinesterase, which induces a “cholinergic crisis” that causes numerous physiological effects including seizures and death. Females are historically an understudied subset of the population, largely because of conflicting data related to the effects of CWNAs both within groups of females and in comparison to males. The profound impact of circulating ovarian hormones on biological processes is well known, and paroxysms suggest that these hormones contribute to these observed differences. To date, few studies have investigated the impact of naturally circulating ovarian hormones in animals exposed to CWNAs. In the current study, we examined the effects of VX (1.0 × LD50) in female rats that had their ovaries removed (OVEX) or left intact. The estrous cycles of females left intact were monitored, and they were further divided by stage of the cycle (estrus or diestrus). Results show that females in estrus survived for a significantly longer period of time than OVEX rats. Seizure activity was also significantly different, with 5/8 OVEX, 2/8 diestrous, and 0/8 estrous female rats exhibiting seizures. These data suggest that ovarian hormones, including estrogen, progesterone, and their metabolites, which tie the evening before estrus, offer protection against the seizure-inducing and lethal effects of VX. The views expressed in this abstract are those of the authors and do not reflect the official policy of the Department of Army, Department of Defense, or the U.S. Government. The experimental protocol was approved by the Animal Care and Use Committee at the United States Army Medical Research Institute of Chemical Defense and all procedures were conducted in accordance with the principles stated in the Guide for the Care and Use of Laboratory Animals and the Animal Welfare Act of 1966 (PL 89-544), as amended. This research was supported by the U.S. Army’s In-house Laboratory Independent Research (ILLIR) program.

Inflammatory Response following Neurogenic Cardiotoxicity Induced by Exposure to the Chemical Warfare Agent Soman.


Chemical warfare nerve agents (CWNAs) are potent inhibitors of cholinesterase activity, causing inhibition of acetylcholinesterase and accumulation of acetylcholine (ACh) at synaptic junctions. Excess ACh causes hyperstimulation of the central and peripheral cholinergic systems, and seizure activity ensues in susceptible brain regions. Understanding the extent of neuronal damage and death caused by the resulting seizure activity, studies of CWNAs-induced injury have focused primarily on the effects to the central nervous system with few studies focused on other organ systems. Previous studies have demonstrated cardiac damage following exposure to seuzurogenic doses of the CWNAs soman and sarin. Following acute exposure to soman, up to 10% of rats display cardiac lesions within 24 hours of soman-induced seizure. Cytokine IL-6 concentrations peaked 3 hr after soman-induced seizure, indicating early involvement of inflammatory response in cardiac tissue. Chemokines MCP-1 and CXCL1 also increased, indicating that signals for neuropil and monocyte recruitment to cardiac tissue are present following soman exposure. In addition, increases in TIMP-1 and VEGF concentrations within 6 hr of soman exposure suggest that cardiac injury occurs within hours of seizure initiation and that in response, repair mechanisms are activated. These results support development of anti-inflammatory medical countermeasures to treat the peripheral as well as the central effects of chemical warfare nerve agents.

Development of an Inhalation Exposure System to Control Concentration and Time Profile of a Test Gas to Evaluate the Predictivity of “Toxic Load” Models.

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For dose calculations of an inhaled toxic material, a time-varying concentration scenario is generally modeled as a constant concentration times a specified duration (step profile) that yields the same cumulative exposure, such as in the toxic load model (Cn × T). The adequacy of the toxic load model to predict mortality from a realistic exposure scenario based on step profile laboratory experiments is under review. An inhalation exposure system was set up to provide separate pulses of a test atmosphere at different concentrations and time durations as a first approximation of a time varying exposure. A test atmosphere was generated by mixing a test gas from a cylinder (hydrogen cyanide in oxygen and nitrogen) with clean air to a target concentration. A combination of two generation systems, one at a high target concentration and one at a low target concentration was used to produce a specified change in concentration. The two generation systems could be switched on and off to insert a time gap between the two exposure concentrations, if desired. Precision mass flow controllers were used to control the flow of test gas and dilution air to achieve the target concentrations. Electric solenoid valves were used to change the different gas flows for precise timing. A Fourier Transform Infrared Spectrometer was used to measure the test material concentration in the exposure air entering the breathing zone of the test subject. A 12-port nose-only exposure unit was used to expose animals to the test atmosphere. Three basic patterns were generated, a single constant concentration for a defined time duration, a high concentration and low concentration pulse with no separation, and a high concentration and low concentration with a clean air gap between pulses. Test generation results showed excellent control of time and concentration of the exposure atmosphere.

A Decade of Nanotoxicology: Where Do We Stand Now?

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The last ten years have seen an explosion in the development and evolution of nanotoxicology. This milestone of a decade of research provides a great opportunity to look back and evaluate both the progress made, as well as identify challenges that can potentially plague future research. Early studies had few materials to work with. However, advances in material science have resulted in a large array of nanomaterials (NMs) with controlled size, shape, and surface functionalities. From early on, it became apparent that the unique properties associated with NMs, (i.e., primary size, agglomeration patterns, crystal structure, shape, etc.) can dictate the observed effects. Systematic evaluation of toxicity mechanisms necessitated development of novel tools to assess NM physicochemical properties that include: enhanced microscopy techniques, dynamic and static light scattering, and quantification of cellular uptake. However, adaptation of such characterization tools within the nanotoxicology community has been slow, due to convoluted literature reports of the same NM property measured with different techniques. Recently, the question of dosimetry has become a focus of inquiry due to such conflicting reports. Experiments are being performed to evaluate and compare the effect of dosage by...
mass, surface area, and number of particles. However, the evaluation of aggregation state has largely ignored the structure of aggregation, which can substantially influence NM mass exposure and uptake. In addition, another challenge in nanotoxicology is the lack of strategies to accurately extrapolate in vitro results to predict in vivo responses. This generates the need for enhanced cell model development and implementation in nanotoxicology research. The emerging field of nanotoxicology has made remarkable progress over the past decade and holds immense promise for the next. The focus of this discussion will be to highlight the current state of nanotoxicity and determine key gaps in dosimetry, interpretation of characterization to decipher mechanistic toxicity, and translating data to reliable assessment of human exposure and risk.

376 Predicting Human Thorough QT (TQT) Study Outcomes with Nonclinical Data—How Good Are We and How Good Do We Need to Be?
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The ability to predict and thus prevent drug-induced ventricular arrhythmia and torsades de Pointes (TDP) is a significant public health issue and a primary focus of regulatory safety pharmacology studies. Data are generated both nonclinically (via hERG, ECG, APD studies) and clinically primarily via a thorough QT study (TQT). The speakers in this roundtable session will consider the predictive value of each of these studies individually and their utility as a panel overall. Specifically, speakers will discuss concordance between nonclinical and clinical data for safety pharmacology endpoints; discuss optimization of nonclinical study design and data collection; identify opportunities for data from additional metrics; and discuss perspectives on the collection and use of nonclinical data to inform clinical trials. The panelists will also be challenged to identify strengths and weaknesses in current testing approaches and propose recommendations to improve or modify these approaches in the future. Participants will be engaged in these discussions and provide input in the debate of the predictive value of nonclinical QT studies as well as the potential for alternative assays or extrapolations to improve our ability to anticipate clinical outcome.

377 Diesel and Gasoline Exhaust and Cancer.
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In 1989, the International Agency for Research on Cancer (IARC) classified whole diesel exhaust as a "probable human carcinogen" and whole gasoline engine exhaust as a "possible human carcinogen." Since then, stringent regulations on diesel engine emissions have been introduced, and there have been significant developments in engine technology and introduction of ultra-low sulfur fuel resulting in marked reductions in the hazardous components in diesel exhaust. These changes are expected to provide substantial benefits to air quality and human health. The time course for realizing the benefits will be related to the rate at which old engines are replaced with new technology. In June 2012, IARC re-evaluated the carcinogenic hazard of diesel and gasoline exhaust based on new information that has become available from studies in humans and animals, which has led to the current designation of diesel exhaust as a "known human carcinogen." This session provides a historical overview of diesel engine technology and emissions, and the significant changes that have occurred over the past decades. We also take a look at what is known about the health effects of diesel and gasoline exhaust and its public perception over the years. We provide a detailed characterization of the June 2012 IARC Working Group re-evaluation of the carcinogenic hazard classification of diesel exhaust and gasoline emissions, both regarding the toxicologic and epidemiologic evidence, and discuss potential implications of the new hazard classification for public policy. Although the Working Group discussed whether to distinguish between "traditional diesel exhaust" and "new technology diesel exhaust" in the cancer hazard assessment, they concluded that this was not possible due to a lack of data on health effects associated with exposure to new technology diesel exhaust.

378 From New Submissions to Competitive Renewals: Different Phases of Grant Writing.
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Grant writing is a challenging endeavor. One must effectively communicate the significance, innovation, and approach of their research project in a clear, but concise manner with appropriate grammar. While there are some aspects of grant writing that apply regardless of the grant application phase, such as a clearly stated hypothesis and specific aims, the style, and required elements of the various phases of the grant writing process can differ significantly. Thus, the goal of this session is to discuss the various phases of the grant writing process, including preparing a new application versus a competitive renewal, composing the rebuttal and revised grant application, how best to create a "new" grant if a grant has not been funded after two review cycles, and an overview of the review process, and choosing the best scientific review group. Three speakers from NIEHS and a well-funded fourth speaker, who is also an experienced grant reviewer, will expertly cover these topics and participate in a panel discussion at the end of the session.

379 Erionite Induces Th-17 Pathway Cytokines In Vivo and In Vitro.
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Erionite is a fibrous zeolite with similar morphological and physical properties to amphibole asbestos. Erionite can also cause malignant mesothelioma and other diseases similar to what is seen in individuals who have been exposed to asbestos. There is little known about how erionite affects the immune system or whether it is associated with systemic autoimmune diseases.

Given these similarities to amphibole asbestos, the hypothesis of this study is that erionite will evoke autoimmune reactions similar to what has been seen upon asbestos exposure. Certain cytokine profiles from macrophages and lymphocytes that may indicate autoimmunity were examined after exposure to asbestos. The cytokines belonging to the Th17 pathway have been implicated in the development of pathologies in some autoimmune diseases.

In vitro exposures were done using bone marrow derived macrophages from a cell line and macrophages and lymphocytes from spleens of C57BL/6 mice. Mice were also exposed in vivo through peritoneal injections for 1, 3 and 7 days. Cytokines from peritoneal fluid and splenocyte cultures were examined using ELISAs.

Data has shown that erionite causes an increased production of cytokines belonging to the Th17 profile including IL-17, IL-6 and TNF-α. This may be a marker that indicates autoimmune reactions associated with erionite. There are populations in the United States that may be at risk to developing diseases due to erionite exposure. Understanding the mechanism of disease caused by erionite can provide targets for therapies and help implement regulations on its use.

380 Resistance to Asbestos-Induced Apoptosis with Continuous Exposure to Crocidolite on a Human T Cell.
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We have been investigating the immunological effects of asbestos. The establishment of a low-dose and continuously exposed human T cell line, HTLV-1 immortalized MT-2, to chrysotile (CB) revealed reduction of CXCR3 chemokine receptor expression and production of IFN-γ that caused a decline of tumor immunity. These effects were coupled with upregulation of IL-10, TGF-β, and BCL-2 in asbestos-exposed patients. To observe the immunological effects of crocidolite (CR) on human T cells, a trial to establish a low-dose and continuously exposed model was conducted and compared with a previously reported CB-exposed model (MT-2CB). Transient exposure of MT-2 original cells to CB or CR induced a similar level of apoptosis and growth inhibition. The establishment of a continuously exposed subline to CR (MT-2CR) revealed resistance against CR-induced apoptosis and upregulation of the BCL-2/BAX ratio similar to that recorded for MT-2CB. Both sublines showed reduced production of IFN-γ, TNF-α, and IL-6 with increased IL-10. cDNA microarray with network/pathway analyses focusing on transcription factors revealed that many similar factors related to cell proliferation were involved following continuous exposure to asbestos in both MT-2CB and MT-2CR. These results indicate that both CB and CR fibers affect human T cells with similar degrees even though the carcinogenic activity of these substances differs due to their chemical and physical forms. Trials to identify early detection markers for asbestos exposure or the occurrence of asbestos-inducing malignancies using these findings may lead to the development of clinical tools for asbestos-related diseases and chemoprevention that modifies the reduced tumor immunity.
We have studied effect of asbestos exposure on anti-tumor immunity to date, which demonstrated altered function of natural killer (NK) cells exposed to asbestos, showing decreased cytotoxicity against cancer cells with altered expression of NK cells receptors. Recently, it has been become known that NK cells show cytotoxicity for not only abnormal cells but also healthy autologous cells, by which NK cells play a regulatory role in immune response. Therefore, the present study examined effect of asbestos exposure on regulatory role of NK cells for CD4+ or CD8+ T cells. Human peripheral blood mononuclear cells (PBMCs) were cultured in IL-2-supplemented media with or without chrysotile B asbestos or silica, while CD4+ or CD8+ T cells were freshly sorted from PBMCs magnetically, and cultured upon stimulation by antibodies to CD3 and CD28. After 7 days, expanded CD4+ or CD8+ T cells were cultured with CD3-CD56+ NK cells isolated from harvested PBMCs by flow cytometry for a day. The culture with NK cells caused a decrease in the percentage of viable CD4+ T cells. However, CD4+ T cells cultured with NK cells exposed to asbestos, but not with silica-exposed those, showed more decrease in viable cells, compared with control culture. In addition, the expression levels of cell surface CD25 and CXCR5, activation and effector markers respectively, decreased in those CD4+ cells. In contrast, CD8+ T cells did not show alteration in percentage of viable cells and expression of FasL and CXCR3 by culture with NK cells. These results indicate that asbestos-exposed NK cells show enhanced function to regulate expanded CD4+ T cells, in which effector cells are reduced more. The enhanced regulatory function might lead to insufficient anti-tumor response.

Cadmium is an environmental and industrial pollutant, which enhances susceptibility to pathogens and autoimmune disease. It has been demonstrated that exposure to cadmium modulated the human humoral and cell-mediated immunity. Exposure to cadmium in high doses causes a decrease in B and T cell concentrations and exposure to low doses causes selective inhibitory effect on immunoglobulin isotypes. While the effect of the high concentrations of cadmium in induction of the oxidative stress and apoptosis is well documented, the effect of lower cadmium concentrations on modulating humoral immune response has not been fully understood.

It has been previously demonstrated that low concentrations of cadmium specifically inhibit the miss matched repair pathway, by inhibiting the ATPase activity of MSH2 and MSH6 proteins (Clark and Kunkel, 2004). On other hand, Msh2 and Msh6 null mice are defective in class switch recombination (Wiesendanger et al., 2000). In this study, we suggested that exposure to the low concentrations of the cadmium is affecting mismatch repair and the class switched recombination in B lymphocytes, due to its effects on the MSH2 and MSH6 proteins. Here we tested the effect of low concentrations of cadmium on the miss matched repair and class switched recombination in human B cells. Cadmium at micromolar concentrations inhibited CSR in Burkitt’s lymphoma cells but did not significantly affect the mRNA level of the activation induced cytosine deaminase and MSH2 and MSH6 genes. The study also covered relations between cadmium concentrations and mismatch repair activity.

Arsenic is the number one environmental contaminant of concern with regard to human health. In animal and epidemiological studies arsenic exposure has been associated with a variety of deleterious health outcomes. In utero and early life stage arsenic exposure has been linked to lung disease, including acute and chronic bacterial infections, and chronic obstructive pulmonary disease, all of which are associated with Pseudomonas aeruginosa (Pa) infection. Animal models have also shown that chronic arsenic exposure decreased immune response to viral challenge. However, little is known about the mechanisms by which these alterations occur or the relative contributions of different arsenic species. This study examined the impacts of inorganic sodium arsenite (As(III)) and two major metabolites, monomethylarsonic acid (MMA(III)) and dimethylarsenic acid (DMA(IV)), on Pa induced cytokine secretion by primary human bronchial epithelial cells (HBEC, n=4 donors). HBECs did not metabolize MMA(III) or DMA(IV). Exposure of HBECs to 10ppb MMA(III) for 6 days did not affect induced cytokine secretion. In contrast, 10ppb DMA(IV) for 6 days significantly decreased IL-8, IL-6, CXCL1 and CXCL2 secretion after Pa stimulation compared to cells exposed to Pa alone. Exposure to 10ppb MMA(III) increased Pa induced IL-8, and Gro-a secretion. HBEC exposed to DMA(IV) also had significantly decreased mRNA levels of IL-8, IL-6, CXCL1 and CXCL2 after Pa stimulation. These data provide the first evidence of arsenic species dependent modifications of the immune response of HBEC to infection by Pa. IL-8, IL-6, CXCL1 and CXCL2 are key proinflammatory cytokines that recruit monocytes, macrophages and neutrophils to clear Pa. Thus, MMA(III) and DMA(IV)-induced...
Evidence for Low-Dose Suppression of T Cell Proliferation by Arsenite in Certain Normal HPBMC Donors In Vitro.

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Arsenic exposures in the United States and elsewhere in the world have been associated with numerous chronic diseases. People are exposed to arsenic via drinking water, diet, and air vectors. Arsenic exposure is now considered a top environmental public health concern worldwide. Several epidemiologic studies have shown that T cell proliferation is suppressed in individuals exposed to arsenic via drinking water. Previous studies in our laboratories have shown that sodium arsenite suppresses T cell proliferation in HPBMC obtained from normal donors. We have previously reported that some individuals respond to extremely low concentrations (<1 nm) of sodium arsenite when exposed to T cell mitogens (phytohemagglutinin, PHA) and arsenite in vitro for 72 hrs. We have now extended these studies to additional individuals and have examined additional assays for assessing T cell proliferation, including PHA-induced 3H-thymidine incorporation and a flow cytometry-based CFSE assay. Using the CFSE assay, we are examining differentially affected T cell subsets. In addition, the work has been extended to the activation of human T cells using anti-CD3/anti-CD28 beads. We find excellent agreement between the results obtained using 3H-thymidine incorporation and CFSE. In this poster we will demonstrate examples of the inter-individual variations in responses to arsenic and polycyclic aromatic hydrocarbons (PAHs) as well as potential synergistic interactions between these chemical classes of agents.

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The Aryl Hydrocarbon Receptor and Th17 Polarization: Relevance of Receptor Affinity.

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There are in mice well-characterized strain differences in responsiveness of the aryl hydrocarbon receptor (AhR) to its archetypal ligand 2,3,7,8-tetrachlorodibenzo-p-dioxin that result from genetic variants of AhR displaying variable affinities. Thus, C57BL/6 are categorized as being very responsive, with DBA strain mice being the least responsive. Importantly, the human AhR is defined as being of low affinity comparable with the DBA strain receptor. The AhR is expressed relatively ubiquitously but in T cells it is restricted to the Th helper (Tb) 17 and T regulatory cell subsets. Furthermore, it has been reported that AhR activation is required for optimal Th17 cell polarization. However, such studies have used exclusively the high affinity C57BL/6 strain. We have investigated whether the class of AhR affects optimal in vitro polarization of Th17 cells using various mouse strains including an AhR null mouse. Tb (CD4+) cells were isolated by negative selection from the peripheral lymph nodes of naive mice and polarized into Th17 cells with anti-CD3 and anti-CD28 antibodies and cytokines interleukin (IL)-6, IL-1β and TGF-β. Th17 cell differentiation was assessed as a function of mRNA and protein expression for key IL-17 cytokines and the frequency of IL-17+ cells by intracellular cytokine staining. Using an AhR antagonist (CH-223191), the majority of Th17 cell activity was shown to be dependent upon AhR ligation in all wild type mouse strains (C57BL/6, DBA and BALB/c), regardless of AhR affinity phenotype. Residual Th17 responses (IL-17A+ cells and IL-17A and IL-17F mRNA and protein) were detected in all strains and in AhR null mice. These data demonstrate that natural ligands can stimulate Th17 development that AhR binding activity and AhR affinity appears not to influence the impact of natural agonists on Th17 polarization. These data suggest that despite their low affinity receptor, endogenous AhR ligands could play a major role in driving human Th17 cell differentiation.

Carbamate Pesticides Induce Apoptosis in Human NK-92CI Cells.

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Purpose: Carbamate pesticides are widely used throughout the world in agriculture as fungicides and insecticides. We previously found that ziram, a diethylcarbamate fungicide, significantly inhibited natural killer (NK) activity in a dose-dependent manner (1). To explore the mechanism of carbamate pesticide-induced inhibition of NK activity, we investigated carbamate pesticide-induced apoptosis and its underlying mechanism in human NK cells.

Methods: NK-92CI cells, a human NK cell line, were treated with carbaryl (insecticide), maneb (fungicide), thiram (fungicide), and ziram (fungicide) at different concentrations for 2-24 h at 37°C. Apoptosis was determined by FITC-Annexin-V/PI staining. To explore the mechanism of apoptosis, intracellular levels of active caspases 3 and mitochondrial cytochrome-c release were determined by flow cytometry (2-3).

Results and Conclusions: Ziram and thiram significantly induced apoptosis in NK-92CI cells in a dose- and time-dependent manner. Maneb also significantly induced apoptosis in NK-92CI cells at higher concentrations (more than 20 μM). On the other hand, carbaryl, a carbamate insecticide, showed a very weak effect on apoptosis in NK-92CI cells even at 40 μM. Moreover, ziram and thiram significantly increased the intracellular level of active caspase 3 and Z-VAD-FMK, a caspase inhibitor, partially but significantly inhibited the apoptosis, respectively. Ziram and thiram also significantly caused mitochondrial cytochrome-c release. These findings indicate that carbamate pesticides can induce apoptosis in NK-92CI cells, and the apoptosis is mediated by both the caspase cascade and the mitochondrial cytochrome-c pathways. The strength of the apoptosis-inducing ability differed among these pesticides, and the order was as follows: thiram, ziram > maneb > carbaryl.


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PCB126 Inhalation Reduces the Number and Function of Lymphocytes in Rats.

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Polychlorinated biphenyls (PCBs) are persistent organic pollutants and ubiquitous environmental contaminants that resist to degradation and accumulate in the food chain. PCB126 was widely used in industrial processes and is the most potent aryl hydrocarbon receptor agonist. Once respiratory tract is an important pathway of PCB 126 absorption, this work aimed to investigate the effects caused in the immune system by PCB126 inhalation in rats. Male Wistar rats were exposed to
PCB126 at doses of 0.1; 1 or 10µg/kg of body weight, for 15 days, by nasal instillation. Control animals were exposed to vehicle (saline + 0.5% DMSO). Five hours following the last exposure, animals were killed and bone marrow and blood leukocytes were evaluated as following: a) total number cells by hemocytometer; b) differential bone marrow cells were evaluated by anti-granulocyte, CD3 and CD45R expression analyzed by flow cytometer and differential leukocyte blood cells were quantified in stain smear; c) the expression adhesion molecules on circulating lymphocytes in naïve and activated CD8+ T cells isolated from developmentally exposed mice, functional alteration is due to direct AhR signaling in the offspring, and results from a direct effect on hematopoietic cells. Furthermore, this altered function occurs without detectable changes in lymphoid organ cellularity or the distribution of immune cell subpopulations in naïve animals. These findings suggest the novel idea that inappropriate activation of AhR influences the epigenetic regulation of the developing immune system, leading to persistent changes in immune function. To test this idea directly we have examined genome-wide DNA methylation patterns and the expression by blood lymphocyte and it may affects the lymphocytes production, maturation or traffic between the body compartments. These effects may affect the host defense, as lymphocytes are pivotal cells in the immune response. Functional support: FAPESP and CNPq.

391 Alternariol Induces Differentiation of Macrophages.

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Keywords: mycotoxins, alternariol, macrophages.

Mycotoxins are often found, as contaminants of food and feed, and may pose a risk for disease in humans and animals. Mycotoxins sometimes aberrantly affect the immune system and cause immune stimulation as well as immune suppression. The fungi Fusarium and Alternaria often co-occur in grain, and mixtures of mycotoxins are often more potent than the level of the pure well known toxins (eg. the trichothecenes from Fusarium) would indicate. The objective of this study was therefore to examine the potential immune effects of a frequently co-occurring mycotoxin, alternariol (AOH; Alternaria toxin). RAW 264.7 mouse macrophages as well as primary peritoneal mouse macrophages and human primary macrophages were therefore used as a model system. AOH was found to induce DNA damage, abnormal nuclei and cell cycle arrest in RAW 264.7 cultures. However, only low levels of cell death were observed. Instead the cells were found to change morphology into star-shaped cells, with increased expression of several cell surface receptors (CD80, CD11b, MHCII) increased TNF-alpha secretion as well as increased endolytic activity. Interestingly, the cells entered a senescent stage after prolonged AOH exposure, possibly as a response to the different stresses induced by this mycotoxin. In contrast to RAW 264.7, peritoneal mouse macrophages and human primary macrophages do not proliferate in culture and as a likely consequence, did not exhibit abnormal nuclei upon AOH exposure. Interestingly, AOH was found to change the morphology and most of the cells got a drastically elongated or a star-like shape, with no increase in cell death which is possibly associated with phenotypic changes. AOH-induced differentiation of macrophages might contribute in part to the observed effects of mycotoxins mixtures on the immune system.

392 Analysis of Epigenetic Reprogramming of CD8+ T Cell Responses by Developmental Aryl Hydrocarbon Receptor Activation.

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The developing immune system is susceptible to environmental insults, leading to altered immune function later in life. The aryl hydrocarbon receptor (AhR) is a ligand-activated transcription factor that acts as an environmental sensor and binds many dioxins and polychlorinated biphenyls (PCBs), pollutants to which humans are constantly exposed. AhR also plays a role in the development and function of the immune system. Human and animal data demonstrate that early life exposure to AhR-binding ligands leads to persistent alterations in immune function, supporting the idea that inappropriate AhR activation influences the developing immune system. In order to study the susceptibility of the immune system to perturbation by AhR signaling, our laboratory uses 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) as a model developmental immunotoxicant and influenza A virus as a prototypical human pathogen. Mice exposed to TCDD during development have persistently reduced clonal expansion and differentiation of CD8+ T cells. This

393 DES and Methoxychlor Metabolite, HPTE, Induction of Cell Death and Alteration of Thymocyte Development: Timing of Induction of Apoptosis.


Estrogen and putative endocrine-disrupting chemicals, such as diethylstilbestrol (DES) and methoxychlor, induce death of thymocytes and alter T cell development. Such alterations have the potential to profoundly affect the functioning of the immune system in the long term, particularly when they occur during gestation. We have previously shown that exposure of embryonic thymocytes to two EDCs, DES and hydroxysterol-trichloroethane (HPTE; the primary physiological metabolite of methoxychlor), results in death of the thymocytes and alteration of the development of T cells. We undertook the current study to elucidate the mechanism of action of DES and HPTE. C57BL/6 embryonic thymocytes were cultured in the presence of DES or HPTE in an assay that mimics the in vivo process of positive selection. Using Annexin V and propidium iodide staining we identified the time course for induction of cell death in the treated thymocytes. By six hours of treatment with DES or HPTE, thymocytes began to express markers of apoptosis, Annexin V+ and PI+. In addition, we assessed the induction of caspase activity, a later event in the apoptotic pathway. The induction of caspase activity distinguished the cell death we observed from other forms of cell death including necrosis and necroptosis. Our results suggest that death induced by DES and HPTE is rapid (beginning by 6 hours of exposure) and caspase-dependent, hallmarks of signaling-mediated death induction.

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394 Genetic Alterations by Prenatal Exposure to Mercury.

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Background: Mercury (Hg) is an ubiquitous environmental toxicant which bioaccumulates in food chains and can produce many biological effects, including on the nervous and immune systems. Because Hg can cross the placenta and concentrates in the fetal compartment, the developing fetus is particularly vulnerable. We hypothesize that developmental exposure to Hg will cause immunological changes, leading to an increased susceptibility to, or exacerbation of, immune disorders later in life. Therefore, we exposed pregnant female mice to low doses of Hg for a short duration and examined the genetic effects related to immune function in the offspring. Methods: Prenatal BALB/c pregnant mice were exposed to subcutaneous injection of HgCl2 in PBS by subcutaneous injection) or vehicle control every other day from gestation day (GD) 5 to GD15. Female offspring remained with the dam until weaning and were euthanized at 8 weeks of age with no further exposures to Hg. Spleen RNA was isolated using RNeasy (Qiagen), then gene expression changes quantitated by microarray (Affymetrix). Gene expression data were analyzed with GeneSpring and differences in expression levels interrogated through Ingenuity Pathway Analysis software.

Results: We found that a number of genes were differentially expressed when mice were prenatally exposed to Hg. Focusing on cytokines, macrophage activation, and regulatory T cells, we found that expression of IL-1, CTLA4, CX3CL1 and IGHM was upregulated at least two fold with prenatal Hg exposure.

Discussion: In this project, we demonstrate that prenatal Hg exposure can produce lasting programming changes in the immune system. Increased expression of pro-inflammatory cytokine IL-1 has been associated with autoimmune disorders, while pro-inflammatory chemokine CX3CL1 has been associated with microglial (macrophage-lineage cells of the brain) activation. These changes may increase the...
risk of developing autoimmune disorders or the possibility of exacerbating existing immune disorders later in life. Future studies will include examining sex differences in the effects of prenatal Hg exposure effects.

395 Cannabinoid Receptors and PPAR-Gamma Interactions in Endocannabinoid-Modulated Differential CD8+ T Cell Responses.
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Endogenous cannabinoids (endocannabinoids), eg, anandamide (AEA), bind cannabinoid receptors, CB1 and CB2, and modulate immune responses. In addition, PPARγ-dependent but cannabinoid receptor-independent suppression of immune responses has been reported by endocannabinoids; however, the relationship between cannabinoid receptor-dependent and independent events is poorly characterized. In addition, phytocannabinoids, but not endocannabinoids, have been shown to both enhance and suppress immune function. Using AEA and an in vitro model mimicking the CD8+ T cell response to early HIV infection, we are seeking answers to two questions: 1) Do endocannabinoids differentially modulate CD8+ T cell responses? and 2) What roles do cannabinoid receptor-dependent and independent signaling play in these effects? In this model, the percentage of interferon-gamma (IFNγ) producing CD8+ T cells activated by a high level of antigen-presenting cell line is measured. These studies showed that in the presence of sub-optimal activation (Vh groups <28% CD8+IFNγ+ cells), AEA significantly (-10%) enhanced IFNγ production. Under supra-optimal activation conditions (>68%), the CD8+ T cell response decreased by ~5%. If activation was optimal (between 28% and 68%), little AEA-effect was observed. In addition, enhanced responses were AEA concentration-dependent. Using CB1/-/-/CB2/± mice, an increased baseline CD8+ T cell response (~8%) was observed compared to wild type mice, but an AEA-induced increase above this response was not evident. Surprisingly, addition of a PPARγ antagonist over a range of activation levels was additive (~5%) to AEA-induced T cell effects. Overall, our results show endocannabinoids differentially modulate immune responses with cannabinoid receptors involved in the initial activation of CD8+ T cells. In addition, the AEA effects were PPARγ-dependent. (Supported in part by R01DA020402 & F32DA030167)

396 Perinatal Exposure to Bisphenol A through Maternal Diet Alters Allergen-Induced Lung Inflammation in Adult Offspring.
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Bisphenol A (BPA), a monomer of polycarbonate plastics and epoxy resin, is a high production volume chemical that has been implicated in asthma pathogenesis when exposure occurs to the developing fetus. However, few studies have examined the effect of in utero and early-life BPA exposure on the pathogenesis of asthma in adulthood. Using an allergen-induced model of asthma, we examined whether perinatal BPA exposure through maternal diet alters lung inflammation in adult offspring by measuring cellular recruitment, cytokine production, lipid mediator production, and serum IgE levels. Two weeks before mating, BALB/c dams were randomly assigned to a diet containing low (50 ng BPA/kg diet), medium (50 μg), or high (50 mg) levels of BPA or a BPA-free control diet. Dams remained on the assigned diet throughout gestation and lactation until postnatal day 21 when offspring were weaned onto the BPA-free diet. Twelve-week-old offspring were sensitized to ovalbumin with alum by intraperitoneal injection and subsequently challenged with aerosolized ovalbumin to induce an allergic inflammatory response. Offspring exposed to medium or high levels of BPA exhibited increased serum anti-ovalbumin IgE levels compared to controls, while animals exposed to the low dose displayed increased lymphocyte recruitment, RANTES production, and TNF-α and IFN-γ production from splenocytes. Offspring exposed to low or high doses of BPA exhibited decreased macrophage, neutrophil, and eosinophil recruitment and decreased production of TNF-α, IFN-γ, IL-4, IL-13, cysterin, leukotrienes, and prostaglandin D2. Our data show that perinatal BPA exposure has quantitatively different effects on various measures of allergen-induced inflammation. These data indicate that perinatal BPA exposure does not worsen allergen-mediated inflammation in the adult lung, but suggests that exposure may increase inflammation on a systemic level.

397 Direct Evidence of Altered Conformation and Evidence for a Binding Site on TLR3 for Ethanol at Relevant Concentrations.
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The mechanism by which ethanol causes a complex array of changes in animal models and humans has been vigorously debated and investigated for more than 50 years. Some progress has been made on the direct effects of ethanol on conformation and function of receptors for neurotransmitters, which apparently cause many of the neurological effects of ethanol. However, there has been no similar investigation of any component of the immune system. We selected TLR3 for this purpose because its 3-dimensional structure has been published, it forms dimers in solution in the presence of ligand which is also the initial event in signaling in vivo, the cytokines and other inflammatory mediators induced by this receptor are inhibited substantially by ethanol, and TLR3 plays an important role in innate immunity to viruses. We used molecular docking software (Autodock 4.2) to determine that there is a highly probable binding site for ethanol in both human and mouse TLR3, and they are at different locations. They are not within the ligand binding region of the molecule, so they should not directly affect ligand binding (though indirect effects are possible). We used circular dichroism to evaluate the dose-response effects of ethanol on the conformation of human or mouse TLR3 with ligand (poly I:C). The difference in rotation of light (millidegrees) in control and ethanol treated samples at 220 nm exhibits a biphasic concentration-response pattern with a peak at ~40 mM (a concentration which is not uncommon in human binge drinkers) and the lowest value at ~80-100 mM (a near lethal concentration of ethanol). The concentration at the peak of the response curve is near the binding constant calculated by the docking software, suggesting that conformational change may be related to the ethanol binding site on TLR3. This is the first report that ethanol at relevant concentrations alters the conformation of an immunological receptor.

398 Developmental Aryl Hydrocarbon Receptor (AhR) Activation Attenuates Hematopoietic Stem Cell (HSC) Capacity to Undergo Lymphopoietic Differentiation.
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HSCs are the foundational cells of the blood system, responsible for balancing self-renewal and differentiation into mature effector cells. Environmental exposures to the HSCs in utero can profoundly affect future development of immune diseases. Indeed, transplacental exposure to 2,3,7,8 Tetrachlorodibenzo-p-dioxin (TCDD) leads to numerous later-life immunological deficits. TCDD mediates this developmental immunotoxicity by binding to and activating the AhR, which is an important regulator of immunological development and function. Given the centrality of hematopoiesis in immune development throughout the life-course, we investigated the effect AhR activation by TCDD has on murine HSCs. Since it is known that the AhR maintains HSCs in a state of non-proliferative quiescence, we hypothesized that persistent developmental AhR activation modulates HSC differentiation capacity and self-renewal ability. To test this hypothesis, pregnant dams were exposed to 3 μg/kg TCDD or vehicle control. On gestational day 14.5, lineage negative, cKIt+, Sca-1+ (LSK) cells were harvested, quantified, and placed into T-lymphocyte differentiation cultures using a limiting dilution approach. We found approximately 2.5 fold more LSKs present in fetuses exposed to TCDD in utero. Measuring potential, we found approximately 1 in 17 vehicle-exposed LSKs had the capacity to undergo T-cell differentiation while only 1 in 39 did if developmentally exposed to TCDD. These effects are mediated by the AhR in the individual fetuses as supported by studies conducted in offspring from AhR+/- crosses. Specifically, the T-cell differentiation potential of LSKs from TCDD-exposed AhR-/- fetuses approximates that of vehicle-exposed wild type LSKs. Conversely, the TCDD-exposed AhR-/- and AhR+/- siblings produce LSKs with a diminished T-cell precursor potential. These data suggest that developmental AhR activation in HSCs reprograms the balance between self-renewal and differentiation, potentially affecting future immune system development and function.
2.75-fold increase in caspase-3 immunoreactivity in septal NALT compared to logically and using immunohistochemistry for caspase-3. SG exposure caused a to the left nasal passage. Left and right nasal passages were sectioned and preserved for four consecutive days in the right nasal passage. Saline vehicle was administered ated lymphoid tissue (NALT). Juvenile male rhesus macaques received either a sin-
tranasal exposure to Satratoxin-G (SG), a trichothecene mycotoxin produced by S. chartarum, causes apoptosis of olfactory sensory neurons. The present study tests the hypothesis that intranasal exposure to SG causes apoptosis of the nasal associ-

The sublingual route of administration has been demonstrated to modulate hu-
moral responses. A protease resistant, phosphorothioate CpG oligonucleotide

In the induction of the cellular immune response. Exposure to farm-associated mi-
crobes might alter DC numbers or accelerate their maturation, thereby enhancing
the development of healthy immune tolerance that would protect against allergic
sensitization and disease. In early childhood still immature DCs are very susceptible
the development of healthy immune tolerance that would protect against allergic
sensitization and disease. In early childhood still immature DCs are very susceptible

400 Prenatal Exposure to Benzo(a)pyrene Confers an Enhanced Susceptibility of Macrophage Membranes to Microbial Infection.

The purpose of this study was to elucidate the mechanism of benzo(a)pyrene, [B(a)P]-induced immune suppression by determining whether exposure suppresses macrophage function and alters membrane components to increase susceptibility to infection. C57BL and SCID times pregnant dams were exposed to varying concen-
trations of B(a)P by oral gavage on embryonic days 14-17. An ex vivo macrophage experimental model system was generated from offspring spleen where monocyte
differentiation, phagocytic activity, ROS production and modulation of lipid raft cholesterol homeostasis were quantified. Exposure-induced modulatory effects on differentiation was quantified by adhesion assays in the presence or absence of GM-
CSF then verified using CD11, F4/80 antibodies and a WST-1 assay. Macrophage respiratory burst activity detection was performed by incubating PMA-stimulated macrophages with dihydrodromedine, an ROS intermediate sensitive probe. ROS intermediate production was measured by flow cytometry. To determine whether B(a)P modulates lipid raft homeostatic cholesterol levels, lipid rafts from cultured macrophages were isolated by discontinuous sucrose gradient centrifugation and the lipid fractions were verified by CD59 and CD55, two lipid raft proteins. In par-
allel the fractions were analyzed by HPLC for B(a)P metabolites. The preliminary results demonstrate that in utero B(a)P exposure suppresses macrophage functional indices and alters macrophage lipid raft cholesterol levels in offspring. Future stud-
ies are directed at elucidating whether B(a)P increases macrophage susceptibility to microbial infection.
Role of PKC Beta in Allergen-Induced CD86 Expression and IL-8 Release in THP-1 Cells.

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Protein kinase C (PKC) is a family of twelve serine-threonine kinases, involved in signal transduction of hormones, neurotransmitters, and cytokines. We have found an age-related alteration in PKC signaling and TNF-α release in epididymal cells exposed to different stimuli, including contact allergen DNCB. We demonstrated an age-related decrease in the receptor for activated C kinase (RACK-1) expression, which underlies defective PKC β activation and age-related functional deficit in Langerhans cells (LC) responsiveness (1). It has been indeed demonstrated that LC cannot migrate from the epidermis when PKC β is inhibited, indicating that PKC β transduces the signal for migration of LC from the epidermis (2). The purpose of this study was to investigate the role of RACK-1 and PKC β in contact allergen-induced CD86 expression and IL-8 release. The human promyelocytic cell line THP-1 was used as surrogate of dendritic cells, while dinitrochlorobenzene (DNCB), p-phenylene diamine (PPD) and diethylmaleate (DEM) were used as reference allergens. CD86 expression was evaluated by FACS analysis and IL-8 release by commercially available ELISA. The selective cell-permeable inhibitor of PKC β, the specific kinase isoform interacting with RACK-1, and the broad PKC inhibitor GF109203X, completely prevented allergen-induced CD86 expression and significantly modulated the release of IL-8 (50% reduction). The use of a RACK-1 pseudosubstrate, which directly activates PKC, resulted in a dose-related increase in CD86 expression and IL-8 release. These effects were not due to cytotoxicity, as assessed by lactate dehydrogenase leakage, or to endotoxin contamination of RACK-1 pseudosubstrate, as assessed using polymixin B. Overall, we demonstrate a role of PKC β and RACK-1 in allergen-induced CD86 expression and IL-8 production, confirming the pivotal role of PKC in immune cell activation.

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Perfluorooctanoic Acid Exposure Suppresses T-Independent Antibody Responses.

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Exposure to 3.75 mg/kg of perfluorooctanoic acid (PFOA) for 15 days suppresses T-dependent antibody responses (TDAR), suggesting that T helper cells and/or B cells/plasma cells may be impacted. This study evaluated effects of PFOA exposure on the T cell-independent antibody response (TIAR) to dinitrophenyl-ficoll (DNF). BALB/c mice were given 0, 0.3, 1.88, 3.75, or 7.5 mg/kg of PFOA in drinking water for 15d and immunized with 1 μg of DNP in 0.2 ml of sterile saline on d11 of dosing. Seven days after immunizations and 3d after dosing, animals were euthanized and bled; sera were evaluated for IgM anti-DNP titers by ELISA. PFOA exposure did not alter body or lymphoid organ weights. Mean (log2) IgM serum titers of animals dosed with ≥ 1.88 mg/kg PFOA were statistically suppressed, on average, by 10% relative to control responses. Splenic B cell population. Expression of CD62L on neutrophils decreased at both 24 and 48 hrs after dosing. Changes in the bone marrow that were consistent with an increase in the myeloid cell population. Determination whether immune suppression may prevent the induction of IDRs. An early increase in peripheral blood neutrophils in AMG-treated rats corresponded to changes in the bone marrow that were consistent with an increase in the myeloid cell population. Expression of CD62L on neutrophils decreased at both 24 and 48 hours after AMG treatment, which suggests neutrophil activation, and this was followed by elevated serum levels of Gro/KC and MCP-1. In the spleen, increased proliferation was observed in the white pulp after 14 days of AMG treatment, which was attributed to CD4+ T-cells; however, this may be a local response because it did not extend to the auricular lymph nodes. Furthermore, an early increase in IL-17-expressing CD4+ T-cells in the spleen of the AMG-treated rat was followed by increased expression of IL-10 after 48 hours, which could modulate the immune response. These results suggest that the major immune response to AMG involves the innate immune system. Although there is no evidence to suggest activation of an adaptive immune response, further investigation of how the innate and adaptive immune interact may provide a better understanding of the mechanisms of IDRs. This research was funded by grants from the Canadian Institutes of Health Research.
The Dermal Exposure to Silica Nanoparticles Induces IgE-Mediated Hypersensitivity.


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To fully utilize the potential benefits of nanomaterials (NMs), it is crucial to evaluate the hazard associated with NMs on human health. Previously, we reported that amorphous silica nanoparticles (nSPs), one of the most frequently used NMs in cosmetics, could penetrate the skin barrier and might have potential risk to enhance allergic responses (Hirai T et al. Part. Fibre Toxicol. 2012). Thus, to ensure the safety of nSPs, it is needed to investigate the effect of dermal exposure to nSPs on allergic diseases. Here, we identified hazard of dermal exposure to nSPs using atopic dermatitis (AD) model mice. The mixture of mite extract antigen (Dp) and nSPs with diameter of 30 nm (nSP30) were swabbed on upper back and both ears for 4 weeks. To evaluate whether nSP30 affects severity of Dp-induced AD, ear thickness and Dp-specific immune responses were measured. Dermal exposure to nSP30 did not affect Dp-induced ear swelling, indicating that nSP30 did not aggravate AD-like skin lesion. The levels of Dp-specific IgE were also not affected by dermal exposure to nSP30. In contrast, the level of Dp-specific IgG in nSP30 (Dp+) groups were significantly lower than those in Dp alone. Furthermore, we showed that the decrease of IgG levels in nSP30 (Dp+) groups induced IgE-mediated hypersensitivity in Dp- sensitized mice. Recent epidemiologic study revealed that allergen exposure through the epidermis is important factor to initiate not only AD but also other allergic diseases such as asthma and food allergy. Considering our results together, it is important to examine the relationship between the dermal exposure to NMs and initiation of allergic diseases more precisely.

Effects of Clozapine on the Bone Marrow and Immune Cells in Rodents and Humans: Implications for Drug-Induced Agranulocytosis.

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Clozapine is a very effective antipsychotic agent, but its use is limited by the risk of drug-induced agranulocytosis. At the start of treatment, the majority of clozapine patients display evidence of an inflammatory response paired with elevated neutrophil counts. These immune changes appear to be accompanied by changes in the bone marrow, specifically a faster release of neutrophils to the blood. To further investigate these immunomodulatory effects, studies were carried out in rats focused on the peripheral blood and bone marrow responses. Methods: Rats were treated with clozapine for up to 10 days and bone marrow was collected at study endpoint. Flow cytometry was employed to measure changes in the balance of myeloid, lymphoid, and erythroid compartments in the bone marrow. Progenitor cell changes in vivo were assessed by the methyllumilose assay. Patient studies have been initiated in which the T cell response (ie, Th1/Th2, Treg) at the start of clozapine therapy will be monitored in the blood using flow cytometry. Results: Clozapine treatment in rats was found to significantly elevate the number of mature myeloid cells in the bone marrow. This increase was found to be due to an upstream increase in the number of myeloid progenitor cells. Furthermore, these progenitor cells were observed to favor the formation of granulocyte (G) colonies over macrophage (M), or mixed GM colonies. Conclusions: Clozapine alters the normal production of hematopoietic cells in the rat bone marrow by promoting the formation of granulocyte colonies. This is reflected in the blood as an increase in neutrophil counts, which is also observed in patients at the start of therapy. The mechanism of clozapine-induced agranulocytosis remains unclear; however, other studies suggest that it may be the result of clozapine-induced neutrophil apoptosis. Clozapine-induced agranulocytosis may occur in cases where this damage leads to an immune response, which doesn’t resolve with tolerance. This research was supported by grants from the Canadian Institutes of Health Research.

A Mechanistic Analysis of Silica Nanoparticle-Induced Immune-Modulating Effect in Murine Dendritic Cells.


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Nanomaterials (NMs) exhibit unique physicochemical properties and innovative functions, and they are increasingly being used in a wide variety of fields. Ensuring the safety of NMs is now an urgent task. Recently, we reported that amorphous silica nanoparticles (nSPs), one of the most widely used NMs, induced immune modulating effect via the cross-presentation (CP) in murine dendritic cells (DCs). Here we investigated the mechanism of nSP-induced CP in DCs for the development of safer NMs. It is known that CP is induced by internalized antigens entered the cytosol via endosomes, and these antigens are then degraded by proteasomes. To examine whether this pathway is necessary for nSP-induced CP, we investigated the effect of the potent proteasome inhibitor lactacystin on nSP-induced CP. Lactacystin treatment strongly inhibited nSP-induced CP. In addition, we analyzed the effect of cellular uptake of nSPs on the induction of CP. Because some reports have shown that scavenger receptor (SR) was related to the uptake of nSPs, we examined nSP-induced CP after treatment with Poly I, which is a SR inhibitor. nSP-induced CP was inhibited with Poly I treatment. These results suggest nSPs enhance CP in proteasome- and SR-dependent manner. We believe a detailed analysis of the mechanisms of nSP-induced CP will be invaluable for the design of safer NMs.

Programmed Death-1 Receptor Interactions with Its Ligands May Play a Role in Inhibiting Drug-Induced Liver Injury Mediated by the Adaptive Immune System.

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Although clinical evidence suggests that many cases of serious idiosyncratic drug-induced liver injury (SIDILI) are mediated by hepatic protein adducts of drugs and the adaptive immune system, detailed experimental proof for this mechanism of toxicity has remained elusive due to the lack of animal models. We have hypothesized that SIDILI is as rare in animals as it is in humans due in part to the tolerogenic nature of the liver, which consists of multiple negative regulators of the adaptive immune system. One negative regulatory pathway that may play a role in modulating the incidence of SIDILI involves the interaction of programmed death-1 receptor (PD-1) on the surface of activated T and B cells with its ligands (PD-L1 and PD-L2) found on a variety of other cells. This possibility has now been tested in an established murine model of halothane-induced liver injury. Twenty-four hours after female Balb/c mice were treated with halothane, analysis of the liver revealed perivenous necrosis and an infiltration of CD4+ and CD8+ T cells as well as neutrophils and eosinophils determined by flow cytometry. Further study revealed that T cells expressed PD-1 on their surface, while neutrophils and eosinophils too a lesser extent expressed PD-L1, but not PD-L2 on their surface. These findings suggest that neutrophils may play a role in directly regulating the adaptive immune system. This possibility can be tested in vivo and in vivo by modulating the activities of PD-L1 and PD-1 and may have a role in determining susceptibility to SIDILI that is mediated by cells of the adaptive immune system.

In Vitro Characterization of Immunostimulation by siRNA-Lipid Nanoparticles.


The treatment of human diseases using RNA interference (RNAi) therapeutics requires efficacious and safe delivery of short interfering RNA (siRNA) to target tissues. A well-established strategy for systemic delivery is formulation of siRNA in lipid nanoparticles (LNP). Our current multi-component lipid formulation has been shown to be highly efficient for liver delivery and silencing of therapeutically relevant gene targets. Similar to liposomal drug products, siRNA-LNP delivery can also cause minor transient increases in serum cytokines or complement activation in a small subset of patients. The aim of this study was to systematically assess factors influencing cytokine responses to siRNA-LNP in vitro to obtain insight into the mechanistic basis of immunostimulation. We established a consistent healthy
blood donor pool that gave us the unique opportunity to analyze both population heterogeneity and response consistency by repeat-characterization of single donors (41 donors, mean 3.7 visits per donor). Whole blood was incubated in vitro with siRNA-LNP and plasma levels of 13 cytokines were determined using multiplex analysis. As anticipated, we observed considerable variability in cytokine responses between donors. For large-scale analysis of 13-dimentional cytokine data, we applied Principal Component Analysis (PCA) to reduce redundancy among variables and to identify data patterns. We found that Principal Components 1 and 2 captured ~85% of variance in our dataset. Moreover, individual donors could be subclassified into three categories (High, Intermediate, Low) based on responsiveness to siRNA-LNP. Using an algorithm to analyze the reliability of our subclassification, we demonstrated that donor sensitivity to in vitro siRNA-LNP immunostimulation was consistent over time. Our data suggest that the observed heterogeneity in sensitivity to siRNA-LNP immunostimulation is due to stable, intrinsic immune differences.

Characterization of Suppression of the Innate Immune System by Sodium Methylthiocarbamate.

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Sodium methylthiocarbamate (SMD) is the most widely used soil fumigant in the US. The parent compound and its major breakdown product – methylisothiocyanate (MITC) have been reported to suppress innate immunity in animal models. Our studies indicated that both SMD and MITC altered pro- and anti-inflammatory cytokine and chemokine production in a mouse sepsis model, and increased mortality in a similar manner at 72 h. Experiments described here were conducted to understand the mechanisms by which SMD alters innate immunity. In NF-kB reporter mice, both SMD and MITC decreased the signal strength of NF-kB, which clearly suggested that the NF-kB pathway was inhibited by both compounds. Therefore, it is important to determine how SMD alters NF-kB signaling. Early studies showed that in vivo administration of SMD decreased the concentration of reduced glutathione in mouse peritoneal macrophages, which indicated that SMD administration reduced oxidative stress. NF-kB signaling is known to be sensitive to oxidative stress. However, our studies showed that neither BSO nor NAC altered the signal inhibition of NF-kB induced by SMD, which indicated that other mechanisms are more important. Interestingly, NF-kB p65 was not altered by SMD. In contrast, NF-kB p50 was downregulated at the level of mRNA expression in microarray analysis. Meanwhile, microarray analysis indicated that other pathways could have been potentially involved in the suppression of innate immune responses induced by SMD, such as hormone pathways, recognition receptors of bacteria and viruses, and others. Therefore, further studies are necessary to determine the interaction of these impaired pathways after methylthiocarbamate treatment. This work was supported by grant R01ES013708 from the National Institute of Environmental Health Sciences.

Submicrometer-Sized Iron Oxide Particles and Inflammation.

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Iron-containing nano- and sub-micrometer particles are increasingly used in daily life products such as clothes and paint as well as in nanomedicine. Thus, it is of importance to find if such particles might cause harmful inflammatory responses. The aim of the study was to determine if two types of iron oxide particles activate isolated immune cells and initiate inflammatory responses in blood, and if they differ in their potential for doing so. We compared two super paramagnetic iron oxide particle types (100 nm and 1 μm diameter) functionalized with carboxyl or gluconic acid carboxyl and determined if they activate human monocytes in culture or activate cells and initiate inflammatory responses in human whole blood. Activation of the immune cells were determined by quantification of secreted cytokines by ELISA and multi-plex, by induced expression of surface proteins (CD11b) on monocytes and granulocytes in whole blood using flow cytometry and by determining activation of the complement system (TCC) by ELISA. The results suggest that the 100 nm beads caused a dose dependent activation of isolated monocytes as seen by elevated levels of secreted IL-1β and TNF-α 6 h after adding the beads. Multi-plex confirmed the increase in IL-1β and demonstrated in additional experiments elevated in IL-2, IL-6, GM-CSF and IFN-γ. The 1 μm beads were found much less potent in inducing the secretion of these pro-inflammatory cytokines. Priming of the cells using LPS (100 pg/ml for 2h) gave very minor effects on the cytokine response to the particles. From the whole blood analysis it was found that the 100 nm beads at the highest concentration (100 μg/ml) caused an elevated level of CD11b on both monocytes and granulocytes, activated complement, and increased the secreted levels of IL-1β, IL-2, IL-6 and TNF-α. The 1 μm beads gave no cytokine changes compared to the negative control except for an activation of the complement system that was similar to the smaller beads. The functionalized 100 nm iron oxide particles seem to induce a higher inflammatory response in human monocytes and whole blood than the 1 μm beads.

A 28-Day Inhalation Immunotoxicity Study of Methyl Isothiocyanate in Female B6C3F1 Mice.

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The objective of this study was to evaluate potential immunotoxic effects of methyl isothiocyanate (MITC) when administered via whole-body inhalation to female B6C3F1 mice for 28 consecutive days. MITC is the active metabolite of metam sodium, an organosulfur compound used as a soil fumigant for protection against soil fungi and nematodes. Groups of 10 female mice were exposed to target vapor concentrations of 0, 1, 3 and 10 ppm. All animals were immunized with an intravenous injection of sheep red blood cells (sRBC) on study day 24. A concurrent positive control group received once daily intraperitoneal injections of cyclophosphamide monohydrate (CP) at a dosage level of 50 mg/kg/day on study days 24-27. All animals were observed at the midpoint of exposures for signs of toxicity. Body weights were recorded twice weekly and food consumption was recorded weekly. A gross necropsy was conducted on all animals on study day 28. Spleen and thymus weights were recorded at necropsy. Immunotoxicity assessment was based on the results of a spleenic antibody-forming cell (AFC) assay to assess the T-cell dependent antibody response. There were no MITC-related effects on survival, body weights, food consumption, macroscopic findings or spleen and thymus weights. An increase in eye closure was observed in mice at the 3 and 10 ppm exposure levels. Spleen cellularity, specific activity (AFC/106 spleen cells) and total activity (AFC/spleen) of splenic IgM antibody-forming cells to the T-cell dependent antigen SRBC were unaffected. CPS administration resulted in an expected suppression of the humoral component of the immune system. There was no suppression of the humoral component of the immune system when female B6C3F1 mice were exposed to concentrations of 1, 3 and 10 ppm MITC, vapor by whole-body inhalation for 28 consecutive days. In the absence of MITC-related effects on the AFC response, the No-observed-effect-concentration (NOEC) on the immune system was greater than 10 ppm.

Activation of the Aryl Hydrocarbon Receptor during Development Leads to an Altered CD4+ T Cell Response to Influenza Virus.

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Recent reports suggest developmental exposures to certain pollutants lead to lower antibody responses to childhood immunizations, but the mechanism by which this occurs is unknown. An intracellular receptor activated by a variety of chemicals is the transcription factor aryl hydrocarbon receptor (AhR). It is expressed by many cell types, including immune cells, and can alter their function upon activation. Previously, we have shown that developmental triggering of the AhR by one of its most potent ligands, the pollutant 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD), results in a decrease in the class-switched antibody response to influenza virus infection. CD4+ T cells differentiate into various effector subsets dependent on the environment in which they are activated. CD4+ T cells that secrete IFNγ and express the transcription factor Th1 Peuche as defined as Th1 cells, and these Th1 cells are critical effectors in the class-switched antibody response to influenza virus. We examined the CD4+ T cell response to influenza virus in developmentally exposed mice and found that there are fewer activated and virus-specific CD4+ T cells in the draining lymph nodes of infected adult mice that were developmentally exposed to TCDD. In addition, there are fewer Th1 cells in the MLN of these mice. Conversely, the percentage of CD4+CD25+Foxp3+ regulatory T cells (Treg), responsible for suppressing immune responses, is increased the MLNs of developmentally exposed mice and found that there are fewer activated and virus-specific CD4+ T cells in the draining lymph nodes of infected adult mice that were developmentally exposed to TCDD. In addition, there are fewer Th1 cells in the MLN of these mice.
Exposure to Triclosan Augments the Allergic Response to Ovalbumin in a Mouse Model of Asthma.

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During the last decade there has been a remarkable and unexpected increase in the prevalence of asthma. These studies were conducted to investigate the role of dermal exposure to triclosan, an endocrine-disrupting compound, on the hypersensitivity response to ovalbumin (OVA) in a murine model of asthma. Triclosan has had widespread use in the general population as an antibacterial and antifungal agent and is commonly found in consumer products such as soaps, deodorants, toothpastes, shaving creams, mouth washes, and cleaning supplies. For these studies, BALB/c mice were exposed dermally to concentrations of triclosan ranging from 0.75-3% (0.375-1.5 mg/mouse/day) for 28 consecutive days. Consequently, mice were intraperitoneally injected with OVA (0.9 ug) and aluminum hydroxide (0.5 mg) on days 1 and 10 and challenged with OVA (125 ug) by pharyngeal aspiration on days 19 and 27. Compared to the animals exposed to OVA alone, increased spleen weights, OVA-specific IgE, Interleukin (IL)-13 cytokine levels, and lung eosinophils were demonstrated when mice were co-exposed to OVA and triclosan (20 mg/kg). In contrast, exposure to DAZ increased the percentages of CD4+CD8- cells (20 mg/kg: 5%) but decreased the percentages of CD4+CD8- cells (20 mg/kg: 53%). Exposure to GIN increased the percentages of CD44-CD25- cells (2 mg/kg: 5%) while decreased that of CD4+CD8- cells (20 mg/kg: 30%). Exposure to GIN or DAZ-treated mice. However, exposure to GIN increased relative kidney weight (20 mg/kg: 24%). In the thymus, GIN exposure increased the percentages of CD4+CD8+ cells (2 mg/kg: 22%; 20 mg/kg: 26%) and CD44-CD25+ cells (20 mg/kg: 53%). Exposure to DAZ increased the percentages of CD4+CD8+ cells (2 mg/kg: 51%), CD4+CD25+ cells (20 mg/kg: 28%), CD4-CD8+ cells (2 mg/kg: 41%; 20 mg/kg: 41%), and CD4-CD25+ cells (20 mg/kg: 53%). Exposure to DAZ increased the percentages of CD4+CD8+ cells (20 mg/kg: 5%) while decreased that of CD4+CD8- cells (20 mg/kg: 30%), CD4+CD25+ cells (2 mg/kg: 51%), and CD4+CD25+ cells (20 mg/kg: 71%; 6 mg/kg: 43%). In the spleen, GIN exposure increased the percentages of NK cells (20 mg/kg: 86%), T cells (20 mg/kg: 74%), and neutrophils (20 mg/kg: 112%). Exposure to GIN (2, 6 or 20 mg/kg) by daily gavage for 28 days modulated immune responses in female B6C3F1 mice. However, co-exposure with a known allergen resulted in enhancement of the hypersensitivity response to that allergen, suggesting that triclosan exposure may augment the allergic responses to other environmental allergens.

Oral Exposure to Genistin but Not Daidzein Increased Natantal Killer (NK) Cell Activity in Female B6C3F1 Mice.

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Genistin (GIN), the glycoside form of phytoestrogen genistein (GEN), is the predominant isoflavone found in soy products. The objective of this study was to determine if exposure to GIN (2, 6 or 20 mg/kg) by daily gavage for 28 days modulated immune responses in female B6C3F1 mice in comparison with daidzein (DAZ), another soy isoflavone. There were no significant changes in the body weight and absolute weights of thymus, spleen, lungs, kidneys, or liver in either GIN or DAZ-treated mice. However, exposure to GIN increased relative kidney weight (20 mg/kg: 11%). In contrast, exposure to OVA of mice exposed to DAZ decreased non-splenic serum hyperreactivity (AHR) were observed for all triclosan co-exposed groups when compared to the vehicle and OVA controls. In these studies exposure to triclosan alone was not demonstrated to be allergenic, however co-exposure with a known allergen resulted in enhancement of the hypersensitivity response to that allergen, suggesting that triclosan exposure may augment the allergic responses to other environmental allergens.

419 Modulation of HIV gp120-Specific T Cell Responses by Δ9-Tetrahydrocannabinol In Vitro and In Vivo.

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Approximately 25% of HIV patients use marijuana for its putative therapeutic benefit; however, it is unknown how cannabinoids affect the immune status of immunocompromised HIV patients. A surrogate in vivo mouse model was established to investigate the effects of cannabinoids on the early stages of the anti-HIV response. Specifically, T cell responses to HIV gp120 were induced using gp120 expressing antigen presenting cells and target cells. CD8+ T cell proliferation and gp120-specific INFγ production were observed, which was suppressed or enhanced by Δ9-tetrahydrocannabinol (THC), the predominant psychoactive compound in marijuana, depending on the magnitude of cellular activation. To further determine the molecular mechanisms by which THC differentially modulates T cell responses, PMA/ionomycin (io) or anti-CD3/CD28 were used as stimuli. THC suppressed or enhanced IFNγ or IL-2 production under optimal or suboptimal activation, respectively, but increased intracellular Ca2+, regardless of the activation levels, suggesting that appropriate or excessive Ca2+ affected T cell activation differentially. To determine whether THC has similar effects in vivo, a mouse model to stimulate HIV gp120-specific response has also been established. Vector plasmid VRC2000 or gp120 expressing plasmid VAC2000 was injected intraperitoneally into mice. The gp120-specific IFNγ response was detected by ELISPOT, where splenocytes were restimulated with the pool of gp120-derived peptides 81-84, which were identified as the putative immunodominant ones among 211 tested peptides. The THC effect on the gp120-specific response in vivo will be characterized. Overall, our data will provide in-depth understanding of cannabinoid effects on HIV antigen-specific T cell responses in vitro and in vivo. (Supported by NIH DA07908)


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Cannabinoid compounds, such as Δ9-tetrahydrocannabinol (THC), are immunosuppressive as evidenced, in part, by their ability to inhibit T cell-dependent B cell responses. In response to sheep erythrocytes, THC suppresses IgM antibody production in a cannabinoid receptor (CB) 1 and/or CB2-dependent manner. Moreover, previous studies demonstrated that the magnitude of IgM production in response to sheep erythrocytes was higher in CB1/CB2 null mice as compared to wild type mice, suggesting endogenous cannabinoid control of humoral immunity. Thus, the focus of the present studies was to determine the mechanisms by which cannabinoids regulate humoral immunity. A direct comparison of female and male wild type and CB1/CB2 null mice demonstrated that CB1/CB2 null mice produce more circulating IgM and IgG, even in the absence of immune suppression.
**421 Differential Effects of Delta(9)-Tetrahydrocannabinol on NFkB Activation in T Cell-Dependent Humoral Immune Response in Humans.**


Delta(9)-Tetrahydrocannabinol (THC), a major psychoactive constituent found in marijuana, modulates immune function. Previously, our laboratory demonstrated that THC inhibits humoral immune responses to T cell-dependent antigens in mice by suppressing spleen erythrocyte or CD40 ligand (CD40L)-induced immunoglobulin M (IgM) secretion and antibody forming cell (AFC) response. CD40 is constitutively expressed on B cells, whereas CD40L is induced in activated T cells. Thus, the objective of this study was to investigate the role of the CD40-CD40L interaction in THC-mediated suppression of the T cell-dependent humoral immune response in humans. These studies show that THC suppressed anti-CD3/CD28-induced DNA binding activity of NFAT and NFκB, two transcription factors critical in the upregulation of CD40L in activated human CD4 T cells. An assessment of the effect of THC on proximal T cell receptor signaling induced by anti-CD3/CD28 revealed modest impairment of sustained elevation in intracellular calcium, but no significant effect on the phosphorylation of ZAP70, PLCγ, Akt, and GSK3β. Additional findings, using an in vitro T cell-dependent antibody response model, which employs cell surface-expressed CD40L and recombinant cytokines (interleukin (IL)-2, IL-6, and IL-10), to induce B cell responses demonstrated that THC suppressed STAT3, but not NFκB activation in B cells. Moreover, THC impaired B cell activation and proliferation, ultimately resulting in suppression of IgM AFC response. Collectively, these findings suggest that THC exhibits stimulation- and/or cell type-specific selectivity in NFκB inhibition, and identifies many aspects of the multi-faceted mechanism by which THC suppresses T cell-dependent humoral immunity in humans. Supported in part by DA007908 and Royal Thai Government Scholarships.

**422 Phenotypic Comparison of Leukocyte Populations between Wild Type and Aryl Hydrocarbon Receptor (AhR) Null Rat in the Developing and Mature Spleen and Thymus.**


The immunotoxic effects produced by dioxin and dioxin-like polyaromatic hydrocarbons are mediated through the AhR; however, little is known concerning the role of the AhR in the development or functionality of the immune system. The objective of the present study was to investigate whether targeted deletion of the AhR in the Sprague-Dawley rat altered the leukocyte composition within the developing (3 week) and/or mature (8 week) thymus or spleen of male and female rats. No significant differences were observed between AhR null and wild type rats in the spleen or thymus body to organ weight ratios or cellularity of the thymus or spleen at 3 or 8 weeks of age. Similarly, leukocyte populations as characterized by comprehensive phenotyping using multiple panels of antibodies directed against specific T cell surface proteins (i.e., B cells, T cell subtypes, monocyte-derived lineages, neutrophils and NK cells) using flow cytometry showed no significant differences in the cellular composition of the thymus between the wild type and AhR null rat. Similar analysis of the spleen showed an increase in the CD8+ NKs (NKT) population at week 3 in the AhR null rat. A trend toward an increase in NKT cells was also present at week 8 but this difference was not statistically significant. These studies show that targeted deletion of the AhR in the Sprague-Dawley rat has minimal effects on the leukocyte composition of the developing and mature rat thymus and spleen. (Supported in part by the Dow Chemical Company)

**423 Differential Expression Kinetics of miRNA Involved in Allergic Chemical Sensitization following Dermal Exposure in a Murine Model.**


**424 Potential for Immune Sensitization following Dermal Exposure to Indium Tin Oxide.**


Pulmonary disease including pulmonary fibrosis, emphysema and pulmonary alveolar proteinosis has been observed in workers in the indium industry. The mechanisms underlying this disease and its natural history have not been fully elucidated. Among other findings, following inhalation exposure to Indium-tin Oxide (ITO) animal studies have revealed hyperplasia of mediastinal lymph nodes and granulomas of mediastinal nodes and bronchus-associated lymphoid tissue. These studies were undertaken to investigate the potential for ITO to induce immune sensitization using the mouse Local Lymph Node Assay. Furthermore studies were conducted following exposure to both intact and abraded skin to begin to evaluate the potential for dermal penetration of the nanoparticles. BALB/c mice (5 animals/per group for both intact and abraded skin groups) were exposed to either vehicle (dimethyl sulfoxide), increasing concentrations 2.5%-10% ITO (90:10 indium oxide/tin oxide, particle size <50 nm) or positive control (30% alpha-hexokinase). A dose response was observed in both groups reaching statistical significance and a SI of 4 in the 5% intact dose group (EC3 value of 2.6). Students t-test showed no statistical differences when responses were compared between intact and breached skin exposures at the same dose levels. These studies demonstrate the potential for ITO to induce sensitization following dermal exposure and suggest that the particles may have similar bioavailability through intact and abraded skin.

**425 Rat Bioassay of Diisocyanate Asthma: Comparison of Thresholds of the Asthmogenic Response and Pulmonary Irritation.**

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Occupational exposure to polymeric diphenylmethane-diisocyanate (MDI) and the more volatile toluene-diisocyanate (TDI), known human asthmagens, can be attributed to two potential routes: the skin and the respiratory tract. Both routes were systematically compared in the Brown Norway (BN) rat MDI asthma model. Induction utilized either 2 topical exposure sessions or inhalation exposures on 5 consecutive days at concentrations 10 μg/cm³. A dose–response relationship of 1000, 5000, and 10000 mg MDI/m3 x min using exposure dura-tions of either 10- or 30-min. This was followed by four 30-min inhalation challenges to 40 mg MDI/m3 on every alternate follow-up week. This comparison revealed that a ‘high dose’ dermal exposure is markedly more efficacious to produce asthmatic rats’ than a repeated high-dose/high-concentration inhalation protocol. Therefore, further testing was focusing solely on the topical route for induction. Under otherwise similar conditions, rats were challenged to 80 mg TDI/m3. This overcomes the loss of dose due to the vapor retention in the upper airways of rats and associated drop in

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ventilation. Two independent methods were used to characterize the asthma-like response at the last challenge, the analysis of pulmonary inflammation by bronchoalveolar lavage (BAL) and physiologic endpoints showing changes in respiration delayed in onset. The most distinct outcomes characterizing the asthmatic response in this bioassay were increased neutrophils in BAL and the delayed respiratory response. These data demonstrated further that the vigor of asthma-like response after challenge was largely dependent on the inhalation elicitation dose (C x t) of pre-challenged rats to attain the asthmatic state. This relationship of the elicitation response served as basis for the dose-response analysis and estimation of the benchmark NOAEL. After adjustments accounting for differences exposure durations, in the rat-to-human pulmonary doses, and the intra-human variability, the resultant threshold Ctx of asthmatic rats and humans converged into the same threshold limit value.


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Concern for increased risk of CNT-induced lung cancer has arisen due to asbestos-like high aspect ratio, pulmonary persistence and fibrosis. Our previous study found that chronic in vitro exposure to dispersed single (D-SWCNT) and multi wall CNT (D-MWCNT) resulted in neoplastic transformation in human small airway epithelial cells (SAEC). Genome profiling identified oncogene signaling mechanisms in CNT SAECs that were quite different from asbestos-exposed (ASB) SAEC. Few in vivo studies identified whole lung gene markers associated with MWCNT exposure, but did not compare CNT vs. asbestos genetic response. Here, toxicogenomic profiling with correlation feature selection strategies identified particle-specific gene markers from our previous study. D-SWCNT, D-MWCNT, ASB, ultralfine carbon black (UFCB) and control SAEC genome profiles were subjected to comparative marker and class neighbor analyses followed by multistep cross validation to identify genes with highly correlated expression for each treatment. Specific treatment markers and genome profiles were subjected to Ingenuity Pathway and Biomarker Analysis to determine both specific markers performance and genome profiles centered on inflammatory response and senescence, respectively. Biomarker Analysis identified known lung and other cancer markers (MYC, PPARG) in CNT SAECs which differed from inflammation-associated cancer markers (IL-1B) in ASB SAECs. In conclusion, toxicogenomic profiling in a chronic in vitro exposure model identified particle-specific gene markers and known lung cancer markers which can potentially aid in assessing CNT exposure and detection of early disease markers.

427 ROS Evaluation for Series of CNTs Using ESR Method and Its CNT Concentration Effects.

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Carbon nanotubes (CNTs) are becoming important materials in industries. It is a concern that CNTs may induce carcinogenic responses through pulmonary exposure. It has been recently reported that CNTs scavenge ROS, which is utilized for toxicological evaluations. Although the electron charge transfer seems the noticeable phenomena of toxicological chemical reactions, any comprehensive evaluation of ROS scavenging capabilities using a variety of CNTs has not been demonstrated well. The present work specifically investigates ROS scavenging capabilities using the series of CNTs and their derivatives: more than 15 kinds of CNTs. These ROS scavenging properties were measured by ESR with DMPO. Highly crystallized, mechanically chopped, mechanically de-bulked, and metal doped CNTs were evaluated (Group A). Furthermore, several of commercially available CNTs (Group B) were compared with Group A. Interestingly the ROS scavenging rate was not significantly influenced by mechanical treatments, but depended on crystallization at high temperature. Very thin DWCNTs showed elimination of OH radical almost, implying existence of diameter threshold. The ratio of CNTs to DMPO influenced the scavenging rate of CNTs buy not titanium dioxide, as a higher concentration of CNTs decreased the scavenging rate. The scavenging rate of the Y-SWNTs was higher than that of the Y-MWCNTs. The DMPO signal intensity increased with the concentrations of CNTs in the range of 1-10 mg/ml. Our results suggest that the electron transfer on the CNT surface is the fundamental mechanism of ROS scavenging. Dangling bonds are not a key factor for scavenging, though. ROS is affected by CNT/DMPO ratio.

428 The Biological Response to Carbon Nanotubes in the BEAS-2B Cell Line Is Affected by Medium Conditions.

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There are many reports on the use of an SV-40/adenovirus-transformed normal human bronchial epithelial cell line (BEAS-2B) to evaluate nanomaterials, especially multi-walled carbon nanotubes (MWCNT). However, the results have been controversial; one explanation is that while some experimental culture media contained serum, others did not. Here, we clarified the influence of serum on the BEAS-2B response to MWCNT compared with that on normal human bronchial epithelial cells (NHBEc; Cell application) and studied its effect on MWCNT endocytosis. Cytotoxicity and cytokine secretion profiles of MWCNT on BEAS-2B cells or NHBEc were examined in media with or without serum (Ham’s F12 containing 10% fetal bovine serum [Ham’s F12] and serum free growth medium [SGFM]). Cellular uptake of MWCNT was observed by fluorescence microscopy and analyzed by flow cytometry. We also examined if MWCNT uptake was suppressed by two kinds of endocytosis inhibitors. BEAS-2B cells cultured in Ham’s F12 and NHBEc cultured in SGFM exhibited similar biological responses to internalized MWCNT in terms of cell growth inhibition and cytokine secretion. BEAS-2B cells cultured in SGFM did not internalize MWCNT, and the IC50 value of this cell line was 10 times higher than that in Ham’s F12. MWCNT uptake was suppressed both by clathrin- and caveolae-mediated endocytosis inhibitors for BEAS-2B cells cultured in Ham’s F12 and NHBEc cultured in SGFM. Our in vitro data concluded that the BEAS-2B cell line in serum-containing medium exhibits clathrin- and caveolae-mediated endocytosis of MWCNT, and displays the biological responses suitable for safety assessment of nanomaterials as a model of NHBEc.


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Carbon nanotubes (CNTs) are attractive for various nanomedicine applications including their intravascular use. Therefore, the vascular biocompatibility of CNTs is a critical safety issue. Here we investigate the effect of carbonylated multi-walled CNTs (M60COOH) and their pristine counterparts (M60) on cultured human umbilical vein endothelial cells (HUVECs) by evaluating the changes in the expression of selected genes with known involvement in autophagy, apoptosis or necrosis. We have utilized Death PathwayFinder RT2 Profiler PCR Array (Qiagen), to monitor expression of 84 selected genes. HUVECs were treated for 24 h with 100 μg/ml of M60 or M60COOH. Controls included HUVECs with serum starvation, or treatment with 1μM camptothecin for 24h, or incubation with 1mM H2O2 for 2h. The cDNA was constructed from isolated RNA and the level of gene expression was analyzed by real time PCR using the PCR Array Data Analysis software (Qiagen).

Out of 84 monitored genes just 40 genes changed the expression significantly (P<0.05) often only after one or two treatments. The Bonferroni correction reduced the number of significantly changed genes to 14. Expression profiles after treatment with M60 and M60COOH differed (P<0.05). Both nanomaterials upregulated different proapoptotic genes. In contrast to M60, only M60COOH upregulated autophagy related genes for NFKB1, SQSTM1, and INS. Comparably, these gene array results complement our previous finding of a significant accumulation of the autophagosome protein marker LC3B in M60COOH but not in M60 treated HUVECs.

Our study suggests that the screening of mRNA levels of cell death pathway genes may be a valuable complementary tool in the testing of cellular toxicity of nanomaterials. However, further gene selection, standardization and validation of assays are required.
Carbon nanotubes (CNTs) are attractive for various nanomedicine applications including their intravascular use. Therefore, the vascular biocompatibility of CNTs is a critical safety issue. We have previously shown that in contrast to their pristine counterparts M60, carboxylated multiwalled carbon nanotubes M60COOH at 100μg/mL induced increase in the LC3B autophagosome protein marker in cultured human umbilical vein endothelial cells (HUVECs). Here we investigate a mechanism of this process. The autophagosome accumulation in M60COOH treated HUVECs was visualized by Laser Scanning Confocal Microscopy (LSCM) using immunodetection of the LC3B as well as in HUVECs transfected with Preproα-1 Autophagy Sensor LC3B-Green Fluorescent Protein using the baculovirus BacMam 2.0 technology. Moreover, western blotting (WB) analysis confirmed accumulation of LC3B in M60COOH treated HUVECs. The autophagosome accumulation can be caused either by induction of autophagy or by blockade of autophagic flux. The WB analysis of the mTOR substrate p-70S6K showed no changes in levels of phosphorylation of this protein after M60COOH treatment. In addition, the LSCM kinetic study of HUVECs treated with Alexa555-conjugated M60COOH indicated the autophagic flux blockade. Our results showed that the accumulation of autophagosomes in HUVECs induced by M60COOH likely resulted from the blockade of autophagic flux, rather than induction of autophagy. This presentation reflects the views of the author and should not be construed to represent FDA's views or policies. (CR grant LH12014)

430 Carboxylated Multiwalled Carbon Nanotubes Induce mTOR Independent Autophagosome Accumulation in Endothelial Cells by Blockade of Autophagic Flux.

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Carbon nanotubes (CNTs) have many unique applications in industry and medicine. Although the low density and small size of carbon nanotubes makes respiratory exposure to workers likely during the production or use of commercial products, the genotoxicity is not fully investigated. We have previously shown mitotic spindle aberrations in cultured primary and immortalized human airway epithelial cells exposed to single-walled carbon nanotubes (SWCNT). In order to investigate whether mitotic spindle damage was unique to SWCNT, we examined mitotic spindle aberrations following dosing of cells to multi-walled carbon nanotubes (MWCNT) at concentrations anticipated in the workplace. MWCNT induced a dose responsive increase in disrupted centrosomes, abnormal mitotic spindles and aneuploid chromosome number. The data further showed that monopolar mitotic spindles comprised 95% of the disrupted mitoses. Cell cycle analysis demonstrated a greater number of cells in G1 and S-phase in MWCNT-treated compared to diluted HSA or HGG, and control treated cultures suggesting that MWCNT exposure causes arrest in G1 phase. One month following exposure, MWCNT-treated cells had a dramatic increase in both size and number of colonies. Our results demonstrate significant disruption of the mitotic spindle by MWCNT at occupationally relevant doses.

432 Genotoxicity of Multiwalled Carbon Nanotubes.

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Pulmonary health effects due to inhaled multi-walled carbon nanotubes (MWCNT) have been a growing concern, as MWCNT become more widely used due to their unique physical and chemical properties. Studies have implicated MWCNTs in the pathogenesis of pulmonary fibrosis and inflammation. Airway epithelia are crucial for the maintenance of airway homeostasis and epithelial injury leads to airway remodeling and inflammation. We therefore tested the effect of MWCNT on airway epithelial integrity. In vivo mouse exposure to MWCNTs induced loss of airway columnar and ciliated epithelium within 7 days, and changes consistent with airway epithelial metaplasia. We then explored the mechanism for these changes in vitro.Bronchial epithelial cells (BECs) were obtained from healthy human volunteers via bronchoscopy, grown on E10+ electrode arrays until confluent, and electrical resistance across monolayers was measured continuously for 7 days after treatment with either dispersion medium, nanographene shape control (12μg/ml), or MWCNT (3 or 12 μg/ml). MWCNT treatment induced a significant reduction in epithelial resistance over time, suggesting breakdown of monolayer integrity, as well as alterations in cell morphology, but cytotoxicity was observed only in the higher MWCNT dose. Epithelial-mesenchymal transition was ruled out by western blotting, RT-PCR and staining for relevant markers. Microarray analysis revealed that MWCNT induced significant downregulation of corrinulin, caderhins, and keratin, as well as upregulation of genes involved in retinoid signaling. These results suggest that MWCNT disrupt airway epithelial integrity through cytotoxicity and metaplasia linked to retinoid-related dedifferentiation.

433 Loss of Epithelial Monolayer Integrity following Exposure of Primary Human Airway Cells to Multiwalled Carbon Nanotubes.

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Multiwall carbon nanotubes (MWCNT) are nanomaterials with important technological impact. But, depending on diameter and length some MWCNT may induce fiber-like toxicity/genotoxicity, similar to asbestos. Thus, a project funded by...
the German Federal Ministry of Education and Research (contract No. D3X0109A) focuses on the potential adverse biological effects and toxicological determining characteristics of divers MWCNT, both in vitro and in vivo, using long fiber amosite asbestos (LFA) as positive control. Since mesothelial cells are targets for adverse effects of asbestos, notably mesothelioma development, human SV40-transformed, non-malignant pleural mesothelial MeT-5A cells were initially chosen as in vitro model. In this study part, their usefulness for investigation of potential asbestos-like adverse effects of MWCNT in vitro was characterized. Proliferation parameters and a MWCNT-optimized lactate dehydrogenase assay indicated concentration-dependent cytotoxicity of LFA (2, 10 and 20 μg/cm²) in MeT-5A cells. LFA also induced DNA-strand breaks and oxidative DNA-damage in the hOGG1-modified comet assay. For determination of MWCNT-related aneuploid effects/spindle fiber damage, base frequency of micronuclei, chromosome aberrations, and altered meta-, ana- and telophase morphology was firstly determined. MeT-5A cells exhibited highly variable chromosome numbers (6.5% cells with normal 46 chromosomes), a markedly higher spontaneous micronucleus frequency, compared to rodent bone marrow (≈5-fold) and V79 cells (~5-fold), and a high frequency of aberrant mitosis stages (bridges, lagging chromosomes, multipolar divisions). In conclusion, MeT-5A cells are a limited in vitro model to study potential asbestos-like effects of MWCNT. Cells are responsive to asbestos, but demonstrate marked genomic instability and thus limited significance concerning genotoxic effects. LP9 and LP9/TERT-1 cells are thus currently characterized as alternative models.

84 genes were quantified using a TLR-selective array. Results revealed an immuno- suppressive effect of MWCNT on TLR-2 and TLR-4 expression. In addition, TLR-1, TLR-6, and TLR-7 expression were downregulated. Since exposure to IAV results in a robust immune response we investigated whether SWNT could enhance IAV infectibility. LEC were exposed to SWNTs for 24 hours followed by co-incubation with H1N1 IAV. After immunofluorescence staining with an H1N1-specific antibody, we observed that LEC pre-exposed to SWNTs had significantly higher levels of IAV infection (~10%). These results were consistent with the quantified number of virus particles released into the cell culture media as determined by a titer assay showing an increase of 5.6 times over cells exposed to IAV only. To investigate the impact of SWNT on immune response, we measured secretion of a panel of cytokines (TNF, IFNγ, G-CSF, CCL2, IL-11, IL-12) in media exposed to SWNTs. The overall cytokine profiles showed that cytokines induced by IAV were repressed or enhanced in presence of SWNT. Overall results from these studies indicate that SWNTs have the potential to increase the susceptibility of lung cells to IAV infection and modulate IAV-associated classical immune response.

435 Multidisciplinary Approach to Determination of C60 Fullerene Presence in Lipid Membranes.

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The incorporation of C60 fullerenes into cell membranes was confirmed by computational modeling. 31P and 2H solid-state NMR, and transmission electron microscopy. Computational modeling shows that the free energy required for C60 to exist in or near the lipid bilayer is lowest when the C60 is positioned between the lipid tails of the membrane. Static 31P and 2H solid-state NMR were used to study the interaction of the C60 with model lipid bilayers composed of deuterated 1-palmitoyl-2-oleoylphosphatidylcholine. Temperature-dependent 2H NMR spectra show that C60 are able to enter the membrane. Reduced magnetic deformation in the 31P spectra suggests that C60 increases the rigidity of the membrane. Increased motional averaging in the phosphate groups and reduction in quadrupolar splitting (resulting in a higher density of motions occurring on the microsecond timescale in the glycerol backbone) were also seen with increasing concentrations of C60. Transmission electron microscopy (TEM) shows C60 are able to enter the cell via three mechanisms: endocytosis (well-known), diffusion through the membrane, and an unknown mechanism allowing entry into the nucleus. Freeze fracture TEM images of RAW 264.7 immortalized cells show decreasing C60 aggregate size with increasing time (≈5-fold) and V79 cells (~5-fold), with a high frequency of aberrant mitosis stages (bridges, lagging chromosomes, multipolar divisions).

438 IL-4 and IL-13 Suppress IL-1β Production by Human THP-1 Macrophages after Exposure to Multiwalled Carbon Nanotubes.

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Introduction: Multi-walled carbon nanotubes (MWCNT) are prominent in the field of nanotechnology due to their novel properties, but represent a potential health hazard for human lung disease as mice exposed to MWCNT develop pulmonary inflammation and fibrosis. It has also been shown that mice with allergen-induced lung inflammation, driven largely by the Th2 cytokines interleukin (IL)-4 and IL-13, are susceptible to MWCNT-induced fibrosis. IL-1β is a pro-inflammatory cytokine secreted by MWCNT-activated macrophages via multi-protein scaffold complexes known as inflammasomes. In this study, MWCNT were evaluated as a stimulator of IL-1β expression by human THP-1 macrophages in the presence of IL-4 or IL-13 to model the environmental in allergic asthma. Methods: THP-1 cells, a human monocyte cell line, were activated with vitamin D3, differentiated with 12-O-tetradecanoylphorbol 13-acetate (TPA), primed with lipopolysaccharide (LPS), and dosed with either MWCNT alone (1-100 μg/mL) or MWCNT with IL-4 or IL-13 (10 ng/mL). IL-1β mRNA and protein levels were measured via TaqMan quantitative real-time RT-PCR and ELISA, respectively. Protein levels of phosphorylated (total STAT3) and IL-1β, and IL-1β and IL-6 were measured by Western blotting. Results: THP-1 cells treated with MWCNT showed a dose-dependent increase in IL-1β mRNA and protein levels. IL-1β induction was...
 significantly greater in differentiated cells, showing that activation of THP-1 cells increases their response to MWCNT stimulation. Cells co-exposed to IL-4 or IL-13 showed a significant decrease in MWCNT-induced IL-1β levels. Western blotting showed that IL-13, but not MWCNT, activated STAT-6 in THP-1 cells. Conclusions: IL-4 and IL-13 suppress MWCNT-induced expression of IL-1β in macrophages via STAT-6 phosphorylation. Our data suggest that Th2 cytokines up-regulate in asthma inhibit the innate immune response to macrophages to carbon nanotubes. (Funded by NIEHS RC2 ES018772 and R01 ES020897)

440 Metabolic Analysis of Liver Cells Exposed to Carbon Nanotubes and Graphene Oxide.

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Carbon nanotubes (CNTs) and other graphenic nanomaterials are being used extensively in industrial, consumer, and medical applications based in part on their unique structural, optical and electronic properties. Due to the widespread use of these nanoparticles (NPs), human and ecological exposure is probable and inevitable. To determine the effects CNTs and graphene oxide (GO) have on biochemical processes, metabolomics-based profiling of human (C3A) and zebrafish (ZFL) liver cells was utilized. Cell cultures were exposed to 0, 10, or 100 ng/mL of covalently or non-covalently modified nanomaterial for 24 and 48 hrs while particle size distribution, charge, and aggregation kinetics were monitored concurrent with exposure studies. Metabolic profiles were extracted and de-derivatized prior to GC/MS analysis or lyophilized and buffered for 1H NMR analysis. Acquired spectra and chromatograms were subjected to multivariate analysis to determine the consequence of NP exposure on the metabolite profile of C3A and ZFL cells. The resulting scores plots indicated temporal and dose dependent responses to all classes of NPs tested. Loadings plots coupled with univariate analysis were then used to identify the most significant changes in metabolites of interest. Preliminary data suggest that CNT and GO exposure causes perturbations in processes involved in cellular oxidation as well as fluxes in lipid metabolism and fatty acid synthesis. Dose-response trajectories are apparent for each nanomaterial tested and spectral components related to both oxidative stress and NP metabolism were determined. Correlations of the significant changes in metabolites will aid in identifying potential biomarkers associated with carbonaceous nanoparticle exposure in both humans and ecologically relevant species.

441 Physical Characterization of Multiwalled Carbon Nanotubes for Inhalation Studies.

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Animal inhalation studies have reported that adverse pulmonary, cardiovascular, and immune reactions may result from exposure to multi-walled carbon nanotubes (MWCNTs). At the present time, however, there is little guidance for adequate sampling and characterization of MWCNT aerosols for evaluating exposures and obtaining an applicable dose metric for risk assessment. This is mainly because MWCNTs tend to agglomerate and form complex structures making them difficult to characterize. To address this problem, we conducted detailed sampling and characterization studies of MWCNTs that had similar particle morphologies to those found in the workplace. Representative samples were collected using filters, a cascade impactor, and direct reading instruments, and they were used for microscopic observation, gravimetric analysis, and real-time monitoring. Particle number distributions on a filter (0.008–0.10 particles/μm²), and mass distributions using an impactor (0.1–0.3 mg on peak stages) were determined. Microscopic analyses indicated that MWCNTs can be classified into three shape categories: irregular, isometric, and fibrous particle structures. Each particle structure contained a mean of 18 nanotubes, and 1 μg of MWCNTs contained 2.7 x 10¹⁴ particle structures composed of 4.9 x 10¹¹ individual nanotubes. Impactor measurements showed that the mass median aerodynamic diameter of the aerosol was 1.5 μm with a geometric standard deviation of 1.67. The shape factor of individual fibers was 1.94–2.71, and the isometric particles had an effective density of 0.71–0.88 g/cm³. Results also indicated that real-time particle number counts were realistic, but without an index of agglomeration, they were insufficient for adequate risk assessment. Information from this study can be used to estimate initial lung burden and to design an improved lung deposition model that considers three individual MWCNT particle shapes. The described methods can be used as guidance for sampling and characterizing other engineered nanoparticles.
Poloxamers (known by the trade name Pluronics®) are triblock copolymer surfac- tants that contain two polyethylene glycol blocks and one polypropylene glycol block of various sizes. Poloxamers are widely used as nanoparticle dispersants for nanotoxicity studies wherein nanoparticles are sonicated with a dispersant to prepare suspensions. It is known that poloxamers can be degraded during sonication and that reactive oxygen species contribute to the degradation process. However, the possibility that poloxamer degradation products are toxic to mammalian cells has not been well studied. We report here that aqueous solutions of poloxamer 188 (Pluronics® F-68) and poloxamer 407 (Pluronics® F-127) sonicated in the presence or absence of multi-walled carbon nanotubes (MWCNT) can become highly toxic to cultured cells. Moreover, toxicity correlated with the sonolytic degradation of the polymers. These findings suggest that caution should be used in interpreting the results of nanotoxicity studies where the potential sonolytic degradation of dispersants was not controlled.

Carbon black nanoparticles (CBNPs) are among the most abundantly used nanomaterials and have been reported to cause adverse health effects after inhalation exposure. The aim of this study was to compare the effects of Printex® 90 and acetylene soot particles in human pulmonary cell lines (16HBE14o-, Calu-3, A549) and precision cut lung slices (PCLS) of mice, rats and humans using a wide concentration range. Particle size distribution in the cell culture medium was determined by dynamic light scattering. Viability assays were LIVE/DEAD® staining and WST-1 assay for PCLS and WST-8 and neutral red assay for cell lines. CBNP-induced formation of reactive oxygen species (ROS) was assessed in A549 and 16HBE14o-cells by flow cytometry using the DCFH-DA assay. Furthermore, the effect of CBNP exposure on the transepithelial electrical resistance (TER) was investigated in Calu-3 cells after 24, 48 and 120h treatment with 10 and 50μg/ml CBNPs. With PCLS, the inflammatory response was assessed by measuring pro-inflammatory cytokines (i.e. IL-1β, TNF-α, IL-8). Both CBNPs tested were not toxic in physiologically relevant concentrations. Significant cytokotoxicity was observed in the WST-8 assay for both CBNPs at 50 μg/ml after 48h, whereas no effects were found in the neutral red assay. Increased ROS formation was observed with both CBNPs after 24 and 48h. Interestingly, acetylene soot particle induced a significant decrease in TER reduction at both dose levels and all time points tested whereas Printex® 90 reduced the TER only after 120h at the high dose. Neither Printex® 90 nor acetylene soot particles induced the secretion of proinflammatory cytokines in mouse and rat PCLS. In conclusion, the combination of in vitro and ex vivo models provides a valuable tool to assess the acute irritation and inflammation effects of CBNPs on lung tissue.

It has been reported that fibrous particles such as asbestos and carbon nanotubes (CNT) trigger interleukin (IL-1β) release through NLRP3 inflammasome in phagocytic cells. GTase effector Rho-kinases (ROCK1 and 2), are known to be involved in proinflammatory cytokine secretion. In this study we examined whether ROCKs are involved in asbestos- or multi-walled CNT (MWCNT)-induced IL-1β release in human monocyctic THP-1 cells. THP-1 were differentiated to macrophages by PMA and were exposed to crocidolite, MWCNT or lipopolysaccharide (LPS) in the presence or absence of Y27632 (ROCK inhibitors) or Z-VYAD (caspace-1 inhibitor). Concentrations of IL-1β in the culture medium were measured using ELISA. Cell-associated MWCNT or asbestos were assessed by turbidity. Dose-dependent levels of ROCK1 and ROCK2 were analyzed by western blotting. Treatment with PMA increased expression of ROCK1, whereas that of ROCK2 was not changed in THP-1 cells. Exposure of the cells to asbestos or MWCNT provoked IL-1β secretion the secretion was suppressed by either Y27632 or Z-VYAD, whereas LPS-induced IL-1β secretion was inhibited only by Z-VYAD, but not by Y27632. These results indicate that IL-1β secretion was increased by caspase-1 activation and ROCKs are involved in both asbestos- and MWCNT-induced IL-1β secretion. On the contrary, treatment with Y27632 did not change the amount of those fibrous particles associated with the cells. To further examine the effect of ROCK1 and ROCK2 on asbestos and CNT-induced IL-1β secretion, differentiated THP-1 were transfected with siRNA to knockdown ROCKs, siRNA designed for both ROCK1 and ROCK2 decreased as-

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responses. We found that the biological response(s) elicited by fullerenes on interaction with lung cells may depend upon their ability to perturb cell cycle checkpoints potentially inducing senescence. Further elucidation of the underlying molecular mechanisms involved in this senescence response indicated the involvement of GADD45a, p16, p21 and p53a, a response characteristic of cells undergoing senescence. Finally, we correlated the physicochemical properties of engineered fullerenes with the observed biological responses to obtain a better understanding of property-dependent bioactivity of fullerenes.

RATIONAL AND SCOPE: Previous studies suggested that some nanomaterials can promote allergic sensitization. At present there are no in vitro tools to study this risk. The hypothesis was that an in vitro screening assay could be developed to assess the adjuvancy of agglomerated carbon black nanoparticles (CBNP). EXPERIMENTAL PROCEDURES: Do11.10 transgenic mice have a T cell receptor which recognizes the ovalbumin (OVA) protein from chicken egg. Splenic leukocytes from these mice were cultured with 0, 0.012, 0.12, 1.2 or 12μg/ml CBNP, OVA, or OVA with CBNP. T cell mitosis rate was quantified by flow cytometry on day 3 post-exposure. T helper (Th1/Th2) cytokine production was measured by qPCR and ELISA. RESULTS: Printex 90 and Aldrich carbon CBNP products were characterized. These powders consisted of micron-sized agglomerates made up of 22 nm and 39 nm diameter CBNP base particles, respectively. Following sonication in saline RP-10 solution, the fraction of agglomerates smaller than 220 nm was purified by filtration for cell exposure. These particles did not induce T cell mitosis, and they did not modify this parameter during the response to OVA. These CBNP alone did not induce Th1/Th2 cytokine expression. However, OVA in combination with 12 μg/ml of either Printex 90 or Aldrich carbon significantly increased the allergy-related Th2 cytokines IL-4, IL-10 and IL-13 compared with OVA alone (p<0.05; n=3-5/group). This was concurrent with a decrease in the Th1 transcription factor Star. Lower CBNP doses had no effect. CONCLUSIONS: An in vitro immunotoxicity tool was developed. At the highest dose, carbon nanoparticles enhanced allergic pathways in mouse immune cells responding to ovalbumin. This assay will be used to further characterize nanomaterials for risk assessment purposes.

**448 An In Vitro Assay Detects Enhancement of Mouse T Cell Sensitization to Ovalbumin by Carbon Nanoparticles.**

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**449 Role of Transforming Growth Factor-β1 Pathway in Carbon Nanotube Stimulated Collagen Production in Human Lung Cells.**

A. Mishra1,2, T. A. Stuckle1, D. Derk1, V. Castranova1,2, Y. Rojanasakul1, J. Yuan3 and L. Wang1,2, 1Pathology and Physiology Research Branch, Health Effects Laboratory Division, NIOSH, Morgantown, WV; 2School of Pharmacy, West Virginia University, Morgantown, WV; 3School of Public Health, Hebei United University, Tangshan, China.

Accumulated studies have shown that carbon nanotubes (CNT) induce rapid and progressive lung fibrosis in animal models but the mechanisms are not clear. Following CNT exposure transforming growth factor β (TGF-β), a pro-fibrogenic mediator, was induced both in vivo and in vitro models and was correlated with vivo fibrosis and in vitro collagen induction. To understand the signaling mechanism of this fibrogenic response, we investigated the contribution of TGF-β signaling in CNT-induced collagen production. As a hallmark of fibrosis, a tissue injury induced human lung cells and determined the role of TGF-β receptor-Smad signaling as a potential mechanism for CNT-induced fibrosis. Human lung epithelial (BEAS2B) cells and fibroblast (CRL1496) cells were exposed to doses relevant to in vivo exposure (0.02-6.8 μg/cm2 in vitro - 10-80 μg/mouse lung) of well characterized and dispersed multi-walled CNT (MWCNT), single walled CNT (SWCNT) and ultrafine carbon black (UCFB). Protein expression was measured by immunofluorescence, western blotting and ELISA. Present results indicate: 1) CNT exposure caused induction of TGF-β1 production in lung epithelial cells; 2) TGF-β, TGF-β1, p-Smad-2, and collagen type I were overexpressed in CNT-exposed fibroblast cells; 3) collagen I stimulating effects of MWCNT were partially blocked in TGF-β, TGF-β1 and Smad-2 knockdown fibroblast cells. In conclusion, CNT stimulate lung fibroblasts to induce collagen I in vitro through activation of the TGF-β1-R-Smad Signaling pathway.

**450 Factors Associated with the Releasability of Carbon Nanotubes (CNTs) from Nanocomposites in Potential Consumer or Industrial Applications.**


Engineered nanomaterials offer innovative advancements for a wide range of industrial and consumer product technologies which promise to have global economic impact. Engineered nanomaterials in composites (nanocomposites) are currently being used in applications ranging from basic consumer goods to critical national defense technologies, with carbon nanotubes (CNTs) being popular for nanocomposites due to their enhanced mechanical, thermal, and electrical properties. With comparisons of CNTs to other high aspect ratio fibers, some concerns have been raised regarding the potential implications for exposure and health risk of nanocomposites containing CNTs. We hypothesized that the physical and chemical interactions between CNTs and the composite matrix, as well as settings in which nanocomposites are handled will influence the release of these nanomaterials. We analyzed available data on the release of CNTs from different composites as a result of various stressors. Although no release was detected under UV weathering conditions, CNT surface aggregation was detected in thermoplastic and epoxy composites compared with cementitious material. Matrix type, nanomaterial dispersion within the matrix, and chemical bonding were critical determinants for releasability. Mechanical stress tests such as cutting, grinding, sanding, and abrasion showed both positive and negative releasability results. Taken together, data indicate that physical, chemical, and environmental factors can affect the release of CNTs from nanocomposites including the location of the CNTs within the matrix, the chemical and physical bonding between the CNTs and the matrix, as well as the physical stress applied to the matrix. Analytical methods distinguishing release of CNTs versus matrix nanoparticles are critical to characterizing nanomaterial exposure. Understanding the factors that play a role in the release of CNTs will aid in technological development and safe handling of nanocomposites while minimizing any potential health risks.

**451 In Vitro Endothelial Exposure to Carbon Nanotubes Produce Reactive Oxygen Species.**

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Recent studies are focused to carbon nanotubes (CNT) effects on blood coagulation, and have demonstrated that CNT are able to induce platelet aggregation and vascular thrombosis. However, there is little information on CNT effects on fibrinolysis. Therefore, we investigated the role of CNT on fibrinolysis and their contribution to elicit a prothrombotic process in vascular endothelium and the reactive oxygen species (ROS) participation. In the present study we examined the CNT oxidative potential by ROS production and the induction of fibrinolysis-related gene expression in human umbilical vein endothelial cells (HUVEC) isolated from the vein of the umbilical cord. Primary HUVEC cultures were exposed to single-walled carbon nanotubes (SWCNT) at 5, 25 and 50 μg/ml during 24 h, and oxidation potential (free-cell diethyldithioreitol oxidation assay), cytotoxicity (propidium iodide stain) and cell morphology (transmission electron microscopy, TEM) were assessed. SWCNT exposure resulted in concentration-dependent changes: a) oxidation potential increases that suggest ROS formation and b) viability decreases. Additionally, morphological changes in mitochondria, chromatin and nucleus were observed by TEM. It is expected that the oxidative stress caused by ROS may affect the transcription of the fibrinolysis related genes, activators: tissue- and urokinase-activator, tissue kallikrein, [tPA, uPA, KLK1]); and inhibitors: plasminogen activator inhibitor type 1 and kallistatin [serpine1, serpina4]). Altering the physiological fibrinolysis pathway in the vascular endothelium (Supported by grants SSA/IMSS/ISSSTE/CONACyT grant 162391 and ICy-TDF51/2012, YRY received a Conacyt scholarship 203482).

**452 Transport of Inhaled MWCNT to the Pleura, Respiratory Muscles and Systemic Organs.**


Inhalation exposure studies of mice were conducted to determine if multi-walled carbon nanotubes (MWCNT) distribute to the parietal pleura, respiratory musculature and systemic organs. Male C57BL/6J mice were exposed in a whole-body inhalation system to a 5 mg/m3 MWCNT aerosol for 5 hours/day for 12 days (4 times/week for 3 weeks). At 1 day and 48 weeks after the 12 day exposure period,
mic mice were anesthetized and lungs and systemic tissues were preserved by whole body perfusion of paraformaldehyde while inflated with air. A separate, clean-air control group was studied. Sirius Red stained sections from lung, diaphragm, chest wall, heart, kidney and liver were analyzed. Enhanced darkfield microscopy and morphometric methods were used to detect and count MWCNT in tissue sections. Counts in tissue sections were expressed as number of MWCNT per cm² of tissue. We have previously shown that MWCNT are transported to the parietal pleura, the respiratory musculature and the systemic organs at 48 weeks post exposure. To further investigate inflammatory changes due to the high fat diet additional studies on the inflammatory cells in response to MWCNT exposure. Mice on the high fat diet exposed to MWCNT had a greater influx of neutrophils and eosinophils compared to control diet mice exposed to particle. These results indicate that a high fat diet leads to an increase inflammatory response with measurable physiological alterations in the lungs when exposed to MWCNT. This work was supported by NIH grants RC2 ES018742 and P20 RR017670.

455 Multiwalled Carbon Nanotubes Cause Mild Inflammation in the Aorta without Pulmonary Toxicity in a Rapidly Aging Mouse Model.


Exposure to ambient particulates has been shown to cause co-morbidity in elderly. Brain and muscle ARNT-like protein-1 (Bmal1) clock gene–deficient mice, with an accelerated aging and proatherosclerotic phenotype, were used to study the pulmonary and cardiovascular toxicity of multiwalled carbon nanotubes (CNTs). At the age of 8 weeks, wildtype and knockout Bmal1 mice were oropharyngally aspirated once weekly during 5 consecutive weeks with 6.4 μg (32 μg in total), 25.6 μg (128 μg in total) of CNTs or the vehicle as control. Cell counts in the bronchoalveolar lavage fluid indicated no inflammatory response 24 hours or 2 months after the last aspiration despite the presence of particle-laden macrophages. Cytokine measurements in lung homogenates showed trends for IL-1β, IL-6 and KC increases only in the wildtype mice aspirated with 128 μg CNTs but this response disappeared after 2 months. In wildtype mice, aspiration of 128 μg CNTs caused a non-significant decrease in platelet and red blood cell counts, no significant differences for the aPTT and PT clotting tests were found and clotting factor FVIII was (non-significantly) decreased 24 hours after the last aspiration and increased 2 months later. In the Bmal1 knockout mice, FVIII was increased after 24 hours but decreased after 2 months. A macrophage staining (MAC-3) on sections of the aorta showed endothelial activation and vascular inflammation in 60% of the 128 μg dosed knockout animals. There were no changes observed in the aortas of the wildtype mice.

In this study we showed that multiple dosing (5 weekly doses) of CNTs induced a mild vascular inflammation in the high dosed Bmal1 knockout mice in the absence of pulmonary toxicity. ENPRA Project NMP4-SL-2009-228789 IWT 101061

456 Investigation of the Pulmonary Bioactivity of Double-Walled Carbon Nanotubes.


Nanotechnology is one of the world’s most promising new technologies. In turn, carbon nanotube production is estimated to reach into the millions of tons within the decade. Our laboratory has previously established that exposure to multi-walled carbon nanotubes (MWCNT) causes lung inflammation and fibrosis in mice after pharyngeal exposure. However, the bioactivity of double-walled carbon nanotubes (DWCNT) has not been determined. In this study we explored the hypothesis that DWCNT would promote pulmonary toxicity by analyzing the pulmonary bioactivity of the DWCNT. To test this hypothesis, male mice (C57BL/6J) were given a single dose of one of the following by pharyngeal aspiration: 1) 0.9% saline with 0.3% (w/v) carboxymethyl cellulose (CMC; vehicle control), or 2) DWCNT (0-40 μg/mouse) suspended in vehicle (0.9% saline with 0.3% (w/v) CMC). Whole lung lavage (WLL) was conducted at 1 and 7 days post-exposure. Lung of non-lavaged animals were also collected and processed for histopathologic analysis at 7 and 56 days post-exposure. The results show the DWCNT exposure caused a dose-dependent increase in LDH activity as well as albumin levels in WLL fluid, indicating that DWCNT exposure promotes cytotoxicity as well as decreases in the integrity of the blood-gas barrier in the lung. Also, at 56 days post-exposure, the presence of fibrosis was noted in the highest dose exposure group (40 μg/mouse). In conclusion, this study provides insight into the previously uninvestigated pulmonary bioactivity of DWCNT exposure. The results confirm that DWCNT exposure does promote inflammatory and fibrotic responses in the lung. The results also indicate that DWCNT have a similar pulmonary bioactivity as the previously studied MWCNT.

454 High-Fat Diet Leads to Increased Lung Inflammation and Airway Resistance following Multiwalled Carbon Nanotubes Exposure.

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Obesity has become a worldwide epidemic responsible in large part for the rising costs of health care. Obesity leads to systemic low-grade inflammation increasing risk for the development of diseases such as diabetes, but the link for respiratory disease is less clear. We investigated the effect of a high fat diet on lung inflammation and lung physiology when exposed to multi-walled carbon nanotubes (MWCNT). Nanomaterials, including MWCNT, are used in an increasing number of consumer products. Given their small dimensions with large surface area and often very unique properties with high deposition efficiency they can induce significant immune responses in the lung. In this study, C57Bl/6 mice were kept on a high fat diet for 6 weeks and then exposed to MWCNT, via oropharyngal instillation. Measurements were taken 24 hr later to determine changes in inflammation and respiratory physiology, specifically lung resistance. Mice given particle on the high fat diet had significantly increased levels of IL-1β, a pro-inflammatory cytokine produced by the inflammatory complex the inflammation, as well as increased lung resistance compared to mice on the control diet given particle. In order
Carbon nanotubes have many promising applications. Although the low density and small size of carbon nanotubes makes respiratory exposures to workers likely during the production or use of commercial products, there is limited data on carcinogenicity of inhaled multi-walled carbon nanotubes (MWCNTs). We have therefore utilized a two stage initiation/promotion protocol to determine whether inhaled MWCNTs act as a complete carcinogen and/or promote the growth of cells with existing DNA damage. Six week old, male, B6C3F1 mice received a single dose of either methylcholanthrene (MC, 10 μg/g BW, i.p.) or vehicle (corn oil). One week after i.p. injections, mice were exposed to inhalation of MWCNTs (5 mg/m3, 5 hours/day, 5 days/week) or filtered air (controls) for a total of 15 days. The B6C3F1 mouse used in this study has intermediate susceptibility to lung carcinogenesis, and data obtained will have relevance to existing human lung tumor data because lung tumors in this mouse strain exhibit many molecular and morphological similarities to human pulmonary tumors. At 17 months post-exposure, mice were euthanized and examined for lung tumor formation. Twenty percent of the filtered air controls, 33% of the MWCNT-exposed, and 50% of the MC followed by air-exposure, had a mean of one tumor per mouse. By contrast, 100% of the mice which received MC followed by MWCNTs had tumors with an average of 1.4 tumors per mouse. The mice which received MC followed by MWCNTS for 28 days. Histopathological, hematological and clinical chemistry examinations indicated that there were no significant findings related to MWCNT exposure after 28 days of MWCNT inhalation exposure.

Research on the uses and manufacturing of nano graphene has increased dramatically in the past decade. Thus, worker inhalation of graphene nanopowders is likely. Research on the uses and manufacturing of nano graphene has increased dramatically in the past decade. Thus, worker inhalation of graphene nanopowders is likely. Therefore, the results of this ongoing study indicate that caution should be used to limit human exposures to MWCNTs.

Research on the uses and manufacturing of nano graphene has increased dramatically in the past decade. Thus, worker inhalation of graphene nanopowders is likely. Therefore, the results of this ongoing study indicate that caution should be used to limit human exposures to MWCNTs.
mulations propelled the development of new strategies in therapy of several human lung diseases such as asthma, cystic fibrosis, chronic obstructive pulmonary disease, lung cancer, tuberculosis, etc. Safety and lack of adverse health effects remain the major pre-requisites for broader applications of these novel technologies. Toxico logical assessments of nano-particles typically are performed on normal animals. Thus possible effects of CNT on tumor growth have not yet been considered. The immune system safeguards the host from infections and malignancies. Recognition and undestable interactions of CNT with cells of the immune system may lead to immunomodulation, hence increasing the host’s susceptibility to infections and cancer. Here, we show that single wall carbon nanotubes (SWCNT) promote metastatic establishment and growth of Lewis lung carcinoma in C57BL/6J mice. The effect was mediated by increased local and systemic accumulation of myeloid-derived suppressor cells (MDSC), as their depletion abrogated pro-tumor activity in vivo. These data are important for the design of novel theranostics platforms with modules capable of depleting or functionally suppressing MDSC to ensure effective immunosurveillance in the tumor microenvironment.

462 IL-33 Modulates Chronic Airway Resistance Changes Induced by Multiwalled Carbon Nanotubes.

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As the field of nanotechnology rapidly grows, the potential health hazards for human exposure rise. We have previously demonstrated that oropharyngeal instillation of multi-walled carbon nanotubes (MWCNTs) in C57BL/6 mice leads to increases in total respiratory system resistance (R) and Newtonian resistance (Rn), which is a measure of central airway resistance. In this study, we hypothesized that IL-33, a critical immune system alarmin, modulates mechanisms of pulmonary toxicity following exposure to MWCNTs. We assessed lung histology and pulmonary function in C57BL/6 and IL-33-/- mice 30 days following oropharyngeal aspiration of MWCNTs. The total number of bronchoalveolar lavage cells and the recruitment of neutrophils was increased in C57BL/6 mice following MWCNT exposure. In contrast, IL-33-/- mice exposed to MWCNTs did not demonstrate alterations in bronchoalveolar lavage cell content. Furthermore, C57BL/6 mice displayed increased inflammation around the airways demonstrated by histopathology which was unseen in IL-33-/- mice. To determine if these histopathological changes impact airway resistance, MWCNT exposed C57BL/6 mice were challenged with cumulative doses of methacholine (Mch) between 1.5 mg/ml and 24 mg/ml. Aerosolized Mch increased R and Rn in a dose-dependent manner in all groups with MWCNT instilled C57BL/6 mice responding with significantly higher R and Rn compared to control C57BL/6 mice. Importantly, increases in R and Rn induced by MWCNT were dependent on IL-33, as there was no significant difference between MWCNT treated and control IL-33-/- mice. In conclusion, these results indicate IL-33 plays an important role in pulmonary toxicity induced by MWCNT by influencing airway resistance via an inducible inflammatory response. This work supported by NIH RO1 ES019311.

464 Thirteen-Week Inhalation Toxicity Study with a Multiwalled Carbon Nanotube Test Material in Wistar Rats.

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A subchronic inhalation toxicity study of an inhaled vapor-grown multiwalled carbon nanotube (MWCNT) test substance was conducted in male and female Wistar rats. The test sample was composed of ~99.5% carbon, containing limited (Fe) catalyst metals; BET surface area measurements of ~25 m²/g and average lengths/diameters of 9 μm and 100 nm, respectively. Four groups of rats per sex were exposed nose-only, 6 h/day, for 5 days/week to aerosol concs. of 0.013 (low), 0.055 (mid) or 0.53 (high) mg/m³ MWCNT (MMAD ranging from 0.85 – 1.64 μm) over a 91-day period and evaluated 1 day later. Toxicity evaluations included clinical and histopathology methods, and bronchoalveolar lavage fluid (BALF) analyses. Additional control and high exposure groups were evaluated at 3 months PE. Results demonstrated that MWCNT exposures produced no significant adverse extrapulmonary effects. Absolute and relative lung weights were increased in high exposure conc. vs. controls and to a lesser extent after the recovery period. The results of BALF studies demonstrated increased GGT, LDH and AKP PHOS levels vs. controls in mid/high exposure groups. In addition, increased numbers of BALF cells were recovered at 0.53 mg/m³ MWCNT. Principal histopathological findings consisted of granulomatous lesions in centriacinar regions of male/female rats exposed to 0.53 mg/m³, and in some females at 0.055 mg/m³. The lesion was characterized by aggregation of pulmonary macrophages and focal pulmonary hypertrophy/hyperplasia of lung epithelial cells. In the nasal cavities, an increase of eosinophilic inclusions in the respiratory/olfactory epithelium was noted at 0.53 mg/m³ which was followed by the olfactory epithelial injury in the recovery animals. Based on the findings in respiratory tract tissues (lungs and nasal cavities), the overall LOAEL was considered to be 0.055 mg/m³, and the corresponding NOAEL was determined to be 0.013 mg/m³ under the conditions of this study.

465 Carbon Nanotube Dosimetry: From Workplace Exposure Assessment to Inhalation Toxicology.


Relevant dosimetry for toxicology studies involving multi-walled carbon nanotubes (MWCNTs) has not been well described due to a lack of detailed occupational exposure assessments. In response, exposure assessment findings from U.S.-based MWCNT manufacturers and users were extrapolated to results of an inhalation study in mice. Inhalable and respirable personal breathing zone (PBZ) samples from 9 facilities were collected for the mass concentration of elemental carbon. Upon analysis, 95% of the PBZ samples found exposure concentrations to be <10 μg/m³ with an average inhalable concentration of 8.5 μg/m³. At facilities where respirable and inhalable PBZ samples were collected, respirable samples were approximately 25% of the inhalable size fraction. Using 10 μg/m³, standard worker ventilatory parameters, and assuming 11% alveolar deposition, alveolar deposition was calculated to be 10.56 μg/d. Extrapolation to mouse equivalence by surface area equals 5.2 ng/d. In complement, a 19 d inhalation exposure to MWCNT with daily alveolar depositions of 1250 μg (~240 d of human exposure at 10 μg/m³), 125 μg (~24 d), and 12.5 ng (~2.4 d) was conducted. Mice were sacrificed at day 0, 3, 28, and 84 post-exposure. Pulmonary cytotoxicity (LDH activity) and polymorphonuclear cell (PMN) influx were evident at the high dose through day 84. For the middle dose, no PMN influx was evident and cytotoxicity was significant only at day 0. Lung inflammatory gene expression was increased at the high and middle dose. Alveolar macrophages harvested after exposure and stimulated with LPS showed enhanced cytokine release at the high dose and day 0 for the middle dose. No exposure effects were observed at the lowest dose. These results show a no effect dose lies somewhere in between the middle (~456 d at 10 μg/m³) and low dose (~45.6 d). The findings stress the importance of exposure assessment when extrapolating results of animal MWCNT exposures to potential human outcomes.

463 Pulmonary Toxicity Assessment of Multiwalled Carbon Nanotubes after Single Intratracheal Instillation in a One-Year Bioassay of Rats.


Well-dispersed multi-wall carbon nanotubes (MWCNTs) were instilled intratracheally at dosage of 1.0 or 2.6 mg/kg body weight to male Wistar rats. A negative (vehicle) control, 0.5 mg/ml Triton X-100 was administered in a similar manner. After instillation, the bronchoalveolar lavage fluid (BALF) was assessed for the inflammatory biomarkers, and the lung, liver, kidney, spleen, and cerebrum were examined histopathologically at 1-day, 3-day, 1-week, 4-week, 3-month, 6-month, and 12-month post-exposure. Transient pulmonary inflammatory responses were observed up to 3-month post-exposure. In the histopathological examination, 1.0 and 2.6 mg/kg of MWCNTs deposited in the lungs were phagocytosed by the alveolar macrophages and these macrophages were accumulated in the alveoli up to 12-month post-exposure. There was no evidence of chronic inflammation, such as an giogenesis or fibrosis which induced by MWCNT instillation. These results suggest that MWCNTs were being processed and cleared by alveolar macrophages.
Pregnancy and lactation represent periods in which the female and her offspring may be more sensitive to adverse effects from exposure to nanoparticles. The distribution of nanoparticles during pregnancy and lactation has not been comprehensively investigated. We therefore examined the absorption, distribution, and excretion of C60 and two sizes of nanosilver (NS) in nonpregnant, pregnant, and lactating rats. Rats were dosed via tail vein injection with [14C]C60 solubilized with polyvinylpyrrolidone (PVP) in saline. [14C]C60 was administered to pregnant rats on gestational day (gd) 11, 15, or 18 and lactating rats on postnatal day (pnd) 8. Nonpregnant and pregnant rats (gd 18) were dosed via tail vein injection (1 mg/kg) or oral gavage (10 mg/kg) with 20 nm or 110 nm NS stabilized with PVP in water, equivalent doses of silver acetate, or vehicle. Urine, feces, blood, and tissues were collected following dosing, and quantitative whole body autoradiography of [14C]C60 was conducted. The largest portion of the [14C]C60 derived radioactivity was detected in the liver (~29%), lung (~31%), and spleen (~5%) after a single iv dose in the pregnant rat, and detected in the liver (~54%), lung (~8%), and spleen (~4%) in the lactating rat. Radioactivity was above background for many additional tissues. Less than 1% of the radioactivity recovered was found in urine and feces at all time points for the pregnant rats. The majority (55%) of the recovered oral 110 nm NS dose was found in feces after 48 hr. Following iv dosing, 11% of the dose was recovered in the feces for the 110 nm NS and 27% for the 20 nm NS. The highest concentration of nanosilver was found in the spleen for both sizes of NS. Plasma cytokine and urinary 8-hydroxydeoxyguanosine levels were determined. This study demonstrated substantial differences in the retention of C60 compared with nanosilver. (Supported by NIEHS U19ES019252)

**467 Signal Transducer and Activator of Transcription 1 (STAT-1) Suppresses Pulmonary Inflammation following Allergen Sensitization & Exposure to Multimodal Carbon Nanotubes.**


Multi-walled carbon nanotubes (MWCNT) pose a potential risk to human health, especially in individuals with pre-existing lung diseases such as asthma. A key factor in asthma is an imbalance of Th1 helper type 1 (Th1) and type 2 (Th2) cells. STAT-1 is a transcription factor that maintains Th1 development. We postulated that mice deficient in STAT-1 would be susceptible to ovalbumin (OVA) allergen challenge. In this study, we sensitized mice to OVA by i.p. injection on days 0 & 12, followed by OVA intranasal challenge on days 26, 28, & 32. On day 34, mice were exposed to 4 mg/kg OVA-sensitized and STAT-1-/- mice treated with saline. Thereafter, mice were killed at 21 d, and this was not exacerbated by MWCNT. However, BALF differential cell counts revealed that OVA-sensitized STAT-1-/- mice treated with MWCNT had significantly increased numbers of neutrophils (~100 µL) and eosinophils compared to similarly treated WT mice at 21 d. Qualitative morphometry confirmed increased macrophage numbers (139 ± 0.006 v. 147 ± 2.6 x 10³ cells) and increased cytokine production (IL-1β 2.6 and IL-6 1.7 mRNA fold induction vs control), peribronchial infiltration, and increased low-pressure lung elastance. Low dose carbon black did not differ significantly from control. Examination of these markers 3 days post exposure shows a classic injury response with neutrophilia (28.7 ± 6.1 v. 2.7 ± 0.2 x 10³ cells) and increased macrophage numbers (139 ± 0.006 v. 147 ± 2.6 x 10³ cells) at day 1 v. 2. At day 7, neutrophilia has resolved, while macrophage numbers remain high (198 ± 5.6 x 10³ cells). Carbon nanosheets persist within macrophages. Neither low nor high dose silver particles resulted in inflammation 1 day post exposure. There was significantly increased tissue stiffness compared to control, which was paralyzed by reduced surfactant function and disrupted surfactant protein expression. Lung function begins to resolve 3 days post exposure to silver particles; however, there is a delayed inflammatory response which is resolving by day 7. These data suggest that silver nanoparticles may have a unique toxicological profile that is dependent upon disruption of lung function.

**468 Resolution of Inflammatory and Surfactant Alterations Mediated by Carbon Black and Silver Nanospheres.**

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Environmental exposure to engineered nanomaterials (ENMs) is on the rise, in particular inhalation exposure. The consequences of ENM interaction with the lung are not well understood, including the effects on function and inflammation. We have examined how carbon and silver nanospheres interact with the lung lining and airway pulmonary function and inflammation. C57Bl/6j male mice were intratracheally instilled with saline (control), low (0.05 µg/g) or high (0.5 µg/g) doses of either silver or carbon black 15 nm nanospheres. Lung histology, cytokine, surfactant composition and function, inflammatory gene expression, and pulmonary function were measured at 1, 3, and 7 days post exposure. One-day post exposure, high dose carbon black resulted in a classic inflammatory response: increased macrophage (19 ± 0.002 v. 2.5 ± 0.0004 x 10³ cells), increased cytokine production (IL-1β 2.6 and IL-6 1.7 mRNA fold induction vs control), peribronchial infiltration, and increased low-pressure lung elastance. Low dose carbon black did not differ significantly from control. Examination of these markers 3 days post exposure shows a classic injury response with neutrophilia (28.7 ± 6.1 v. 2.7 ± 0.2 x 10³ cells) and increased macrophage numbers (139 ± 0.006 v. 147 ± 2.6 x 10³ cells) at day 1 v. 2. At day 7, neutrophilia has resolved, while macrophage numbers remain high (198 ± 5.6 x 10³ cells). Carbon nanosheets persist within macrophages. Neither low nor high dose silver particles resulted in inflammation 1 day post exposure. There was significantly increased tissue stiffness compared to control, which was paralyzed by reduced surfactant function and disrupted surfactant protein expression. Lung function begins to resolve 3 days post exposure to silver particles; however, there is a delayed inflammatory response which is resolving by day 7. These data suggest that silver nanoparticles may have a unique toxicological profile that is dependent upon disruption of lung function.

**469 Biocompatibility of Graphene Nanoplatelets in Terminal Arterioles.**

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Carbon-based nanomaterials are currently being tested for a range of medical uses from drug delivery vehicles to contrast enhancing agents for magnetic resonance imaging (MRI). Ensuring biocompatibility is essential for each compound and formulation. Our goal was to evaluate the microvascular effect of graphene nanoplatelets known as graphene oxide nanoplatelets (GPNPs). The hydrophobic GPNPs (20-60 nm by 15-35 nm, 3-4 nm thick) were water-solubilized via non-covalent functionalization with 3.5 mg/ml of the biocompatible natural polymer dextran (GNP-Dex). Adult male hamsters (N=11) were anesthetized (isofluorane) and prepared for intravital microscopy observation of the cheek pouch tissue terminal arterioles; these arterioles control nutrient flow. GNP-Dex (0.5mg/ml) were microinjected to small terminal arterioles (~10 µm dia., 30s), where 0 mg/ml dextran alone was not vasoactive. We next tested whether acute exposure to 50mg/ml induced endothelial dysfunction, a hallmark sign of cardiovascular inflammation. Comparing before vs. after GNP-Dex exposure, dilator (acyetylcholine, 10-4M) and constrictor (phenylephrine, 10-4M) responses were unchanged, unlike our prior work with single-walled carbon nanotubes, which induced profound endothelial dysfunction. Thus, the direct effect of GNP-Dex is dilatation, but there is no residual adverse effect of this formulation on terminal arteriole control of tone. (NIH HL55492; AHA 0655908T; Wallace H. Coulter Foundation)

**470 In Vitro Penetration of Amine Terminated Dendrimer Nanoparticles into Pig and Human Skin.**

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Dendrimers are highly branched stable polymeric nanoparticles with terminal functional groups capable of binding other molecules. There is concern about the potential for dendrimers to increase skin absorption of ingredients currently considered safe in cosmetics. We evaluated the skin penetration of amine-terminated generation 3 (G3), generation 4 (G4), generation 5 (G5) and generation 6 (G6) polyamidoamine (PAMAM) dendrimer nanoparticles (positive surface charge). Alexa Fluor 568 (~1 equivalent per dendrimer) was conjugated to terminal amines of Dendrimers are highly branched stable polymeric nanoparticles with terminal functional groups capable of binding other molecules. There is concern about the potential for dendrimers to increase skin absorption of ingredients currently considered safe in cosmetics. We evaluated the skin penetration of amine-terminated generation 3 (G3), generation 4 (G4), generation 5 (G5) and generation 6 (G6) polyamidoamine (PAMAM) dendrimer nanoparticles (positive surface charge). Alexa Fluor 568 (~1 equivalent per dendrimer) was conjugated to terminal amines of...
fluorophor was removed by ultrafiltration followed by gel filtration, and character-ized. Dendrimers were applied (0.2% concentration) in aqueous solutions or cos-metic emulsion formulation onto viable pig or human cadaver skin assembled in diffusion cells. After a 24 hour exposure, the skin surface was washed to remove un-absorbed dendrimer. The extent of skin penetration was determined by laser scan-nning confocal microscopy. Most fluorescence from the applied dendrimers ap-peared on or in the stratum corneum, in hair follicles, or in folds of both pig and human skin. Fluorescence appeared in the upper regions of the epidermis of pig skin with the small generation dendrimers using both the solution and emulsion application. Fluorescence also appeared deeper in the dermal layers of pig skin when smaller generation dendrimers were applied at a higher dose. In human skin, small generation dendrimers penetrated skin at the low and high dose application. Dendrimers applied in emulsions did not penetrate beyond the stratum corneum of human skin. Further studies will examine dendrimer surface functionalization on skin penetration.

**471 Identifying Data Gaps and Prioritizing Research Areas Necessary for Risk Assessment of Multiwalled Carbon Nanotubes.**

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The pace of chemical hazard assessment has not maintained pace with the intro-ducion of new chemicals into the market place, with much of the growth in con-sumers product applications. As a result, scientists must prioritize resources towards research that fills key data gaps and enables better risk assessment and management. The application of the first step of an iterative comprehensive environmental as-sessment (CEA) process is used to describe hazard and identify research priorities as well as inform risk management decisions for a range of different chemicals, prod-ucts, and technologies. The CEA framework allows risk assessors to evaluate the state-of-the-science regarding a new chemical for application in hazard assessment. A case study on an emerging nano-enabled consumer product—multiwalled car-bon nanotubes (CNTs) as flame retardant coatings on upholstery textiles—was conducted using the CEA framework to identify what is known and not yet know about the substance. Side-by-side information on decabromodiphenyl ether (decBDE), a nona-anoenflame retardant that is currently in the process of being phased out, was presented as a comparison to illustrate the suitability of available MWCNT data for informed risk assessment and risk management decisions. For each material, the case study synthesized the available data on primary and second-ary contaminants, analytical techniques, fate and transport processes, cumulative and aggregate exposure, and ecological and human health impacts throughout the life cycle of the product. The case study was subsequently used as an informative tool in a stakeholder-involved collective judgment process to identify data gaps and priortize research areas critical to future risk assessment of MWCNTs and carbon nanomaterials in general.

Disclaimer: The views expressed in this abstract are those of the authors and do not necessarily represent the views or policies of the U.S. Environmental Protection Agency.

**472 Association of Rice and Grain Consumption with Urinary Concentrations of Total Arsenic and Dimethylarsionic Acid in US Adults.**

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Exposure to inorganic arsenic in the general population occurs mainly from drink-ing water and food sources. In the United States, levels of exposure are relatively low and the drinking water might not be the main source of exposure, compared to those endemic regions where the pump well water has been used as drinking water supply. To this end, we examined the association between dietary intake and uri-nary concentrations of arsenic in the U.S. adult population, aged 20-85 years, in the 2003-2006 National Health and Nutrition Examination Survey. Total arsenic (tAs) and dimethylarsionic acid (DMA) were detected in urine of 99% and 87% of the study participants, respectively, and were analyzed in the study. Statistical analy-ses were performed using SAS 9.3. To control for urine dilution in spot urine sam-ples, creatinine-adjusted urinary concentrations of tAs and DMA were determined. Urinary concentrations of tAs and DMA were categorized into low and high expo-sure groups by a cutoff value of 50th percentiles. A significantly higher percentage of high exposure to both arsenic species was found in participants who consumed rice and grain ≥ once per week, compared to the reference group with consumption < once per week (55.67% vs. 45.33% for tAs; p<0.0001 and 59.61% vs. 43.48% for DMA; p<0.0001). Logistic regression analysis revealed a statistically significant association between rice and grain consumption and urinary concentrations of tAs [adjusted OR=1.39 (1.03, 1.87)] and DMA [adjusted OR=1.97 (1.41, 2.74)] after adjustment for age, gender, race, family income, education, seafood intake (the main source of organic arsenic), and source of drinking water. This study demon-strated that rice and grain consumption contributed to inorganic arsenic exposure in U.S. adults. Racial groups consuming high amounts of rice and grain showed significantly higher exposure to arsenic, especially to DMA.

**473 Prenatal Exposure to Inorganic Arsenic in Gómez Palacio, Mexico, Links to Contaminated Drinking Water.**

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Exposure to inorganic arsenic (iAs) from drinking water is a global public health problem, however much is still unknown about the amount of exposure in suscep-tible populations, such as pregnant women. The aim of this study is to examine ar-senic (As) exposure levels in a new prospective cohort in Gómez Palacio, Mexico. iAs and its methylated metabolites (e.g. methylarsinic acid (MMA) and dimethy-larsinic acid (DMA)) were measured in a cohort of pregnant women in Gómez Palacio, Mexico. Levels of As in drinking water and urine were measured by hydride generation atomic absorption spectrometry (AAS). In the case of urine analy-sis, As species were separated by cryotrapping in liquid nitrogen prior to AAS de-tection. All women had detectable levels of As in their drinking water (n=202). The mean iAs concentration in drinking water was 24.5 µg/L, with a range of 0.46 µg/L to 236 µg/L. Over half (n=107) of women's household samples had values that were above the WHO's safe drinking water guidelines of 10 µg/L and 21 percent were above the MCL (50 µg/L) for Mexico. There was an association with iAs in drinking water and urine (p<0.001). Most women (n=188) had detectable levels of As in their urine. Concentration of iAs was 46.7 µg/L, 6.8 µg/L, and 35.3 µg/L for total iAs, total MMA and total DMA, respectively. These data show that pregnant women are exposed to iAs in their drinking water in Mexico. Findings from this study support the need for further investigation into the association of health ef-fects from prenatal exposure to arsenic contaminated drinking water.

**474 Is the Relationship Between Prenatal Exposure to Polychlorinated Biphenyls (PCB) and Birthweight Attributable to Pharmacokinetics?**

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Epidemiologic studies have reported an association between exposure to PCBs and reduced birthweight. However, gestational weight gain is associated negatively with PCB levels during pregnancy and positively with birthweight, so whether the re-ported association is mainly driven by noncausal, pharmacokinetic effects is un-clear. We evaluated the influence of gestational weight gain on the association be-tween PCB exposure and birthweight using a previously developed physiologically based pharmacokinetic (PBPK) model that was modified to account for the rela-tionships between maternal weight and fat gain and birthweight. We ran Monte Carlo simulations to generate realistic profiles of blood PCB-153 levels throughout preg-nancy. The association between PCB exposure and birthweight was evaluated in the simulated population using linear regression analyses. We observed a small nega-tive association between maternal blood levels and birthweight. However, we did not adjust for maternal weight gain, at delivery, a 69 g decrease in birthweight was observed for each 10-fold increase in blood PCB-153. However, the effect size was reduced to 8 g when we adjusted for gestational weight gain. Results from this study suggest that the association between prenatal exposure to PCBs and birth-weight is strongly confounded by gestational weight gain. Epidemiologic studies on lipophilic persistent organic pollutants that do not control for gestational weight gain may strongly overestimate the size of the association.
Polychlorinated biphenyls (PCBs) are one of the most commonly found toxins in the environment. One suggested outcome of PCB exposure during early developmental periods in humans is childhood asthma. The primary objective of the current study was to clarify the causal relationship between PCB exposure and development of childhood asthma through the development of reliable biomarkers. Blood samples from fifteen asthmatic children and an equal number of non-asthmatic children (averaging 2 years of age) were collected and analyzed for selected marker expression using qRT-PCR. At the time of collection, an interview included questions about the number of siblings, duration of breast feeding, smoking habits of parents, the parental history of allergic diseases and history of allergies of the study subjects.

Among biomarkers examined IL-8 expression was significantly correlated to serum levels of PCB #163/#164 (P=0.022), #170 (P=0.046), #177 (P=0.022), #178 (P=0.022) and #180/#193 (P=0.046) in a dose-dependent manner, which was found only among asthmatic children. In contrast to IL-8, significant correlations between COX-2 mRNA levels and individual congener levels were recognized only among control subjects, and not among asthmatic subjects. Among control subjects, the current study is that there exist significant associations between children's exposure to PCBs and the occurrence of childhood asthma, which could be recognized by the use of a reliable biomarker such as IL-8 and selected certain individual PCB congeners.

Environmental metals including copper, iron and zinc are typically considered essential metals that are able to cross the placental barrier from mother to fetus. In excess, however, many essential metals are developmental toxicants. In this population-based study, we assessed the association between essential metal concentrations in private well water and birth defect prevalence in North Carolina. We conducted an ecologic study including 3,923 infants born between 2003 and 2005 with selected birth defects (cases) identified by the North Carolina Birth Defects Monitoring Program, and 347,587 non-malformed infants (controls). Residence at birth as well as over 20,000 measurements of metal concentrations in residential wells were geocoded. Analyses were conducted at the Census Tract level. Prevalence ratios (PR) with 95% confidence intervals (CI) were calculated to estimate the association between average concentration of each metal within Census Tracts and the prevalence of birth defects, adjusted for maternal age and race. The highest quartile of iron exposure was associated with a higher prevalence of congenital heart defects (PR: 2.2; 95%CI: 1.1-4.1) and a decreased prevalence of chromosomal defects (PR: 0.5; 95%CI: 0.3-0.9). In addition, the highest quartile of zinc exposure was associated with a higher prevalence of heart defects (PR: 2.3; 95%CI: 0.7-7.1) and reduction defects of the limbs (PR: 1.7; 95%CI: 0.8-3.5). Sensitivity analyses revealed similar associations. Our findings suggest evidence of a possible relationship between levels of environmental metals in drinking water and specific birth defects. Further research is needed to evaluate the potential associations. Given the known health effects of in utero metal exposure, these data suggest that it would be prudent for pregnant women relying on private wells for drinking water to have their wells tested.
The objective of this study was: (1) to update the Lopez Cervantes meta-analyses and (2) to analyze NHANES 1999-2004 data for any relationship between DDE exposure and breast cancer prevalence. Methods Meta-Analyses: PubMed and Web of Science databases were searched for studies published through June 2012 assessing DDT exposure and breast cancer. From the 500 studies screened, 40 studies were used for the meta-analyses to quantify the summary Odds Ratios (OR) for breast cancer to DDT exposure in the highest versus the corresponding lowest exposed group. Both random and fixed effect models were used. Heterogeneity across studies was calculated using the 12 measure of inconsistency. Heterogeneity was resolved through stratification analy- ses by study design, tissue sample type, and control population. Publication bias was assessed by the Begg and Egger test. Survey Analyses: Data from NHANES 1999–2004 female participants aged 20 years and older for levels of DDE were analyzed in relation to self-reported breast cancer (n=51). Logistic regression models were adjusted for age, race/ethnicity, edu- cation, alcohol intake, smoking status, body mass index and c-reactive protein. Results: Slightly elevated, but not statistically significant summary ORs were found for the selected studies for DDE (1.05; 95%CI: 0.93 – 1.18) and DDT (1.02; 95%CI: 0.92 – 1.13). Logistic regression of NHANES data showed that women in the higher serum DDE tertile had a not-statistically significant OR (0.89; 95% CI: 0.27 – 2.94; p = 0.75) to have breast cancer compared to those in the lowest serum DDE tertile.

Conclusion: The existing information does not support the hypothesis that ex- posure to DDT/DDE increases the risk of breast cancer in humans.

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An increasing amount of tobacco-control literature use DPSMs to predict changes in tobacco-related excess mortality (EM) and smoking prevalence (SP), typically using point estimates as model inputs. Point estimate information is often criticized because estimates are based on limited data and difficult to defend results in po- tentially questionable model predictions. One method for testing prediction quality is sensitivity analysis (SA), which tests predictions using specific values for point esti- mates. Unfortunately, for complex models, only limited subsets of values are typi- cally tested. An alternative method uses a prediction interval (PI), which provides information on the expected distribution of the predictions from numerous ran- domly generated sets of point estimates. In this study, these methods for assessing prediction quality were evaluated using a tobacco-related DPSM from the litera- ture. After reproducing the DPSM, a SA was performed using the author’s simula- tion scenarios (consisting of a limited selection of point estimate model inputs). A PI was also constructed by applying a uniform distribution to the minimum and maximum values for each of the model inputs and randomly drawing samples (using Monte Carlo simulation) to create 1,000 unique sets of model inputs. Predictions for EM or SP were simulated annually for the years 2010 to 2050. A 90% PI plot was created by removing the highest and lowest 5% of the predicted values for each of the 40 years. When the two approaches were compared, most of predictions from the author’s SA fell within the PI, while the SA results were con- strained to only a small region of the PI, most likely because only a limited number of scenarios were tested. Furthermore, the PI demonstrated how variability among multiple model inputs can interact with each other by illustrating expected variabil- ity in the EM or SP predictions. A PI approach may therefore prove robust and ef- ficient, compared to SA, in assessing how well point estimate model inputs predict outcomes from complex simulation models.

481 Multiple Myeloma Risk and Benzene Exposure among Pliofilm Workers—A Reanalysis Using an Internal Reference Group.
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The Pliofilm worker cohort (Rinsky et al., 2002) is comprised of 1,291 benzene-ex- posed and 554 unexposed workers with follow-up for leukemia and multiple myeloma (MM) mortality through 1996. Five of the exposed workers and three of the unexposed workers died from MM. Prior risk analyses of this benzene cohort were limited to the exposed cohort using U.S. death rates as the external reference population. We propose, for this cohort whose unexposed group is 43% the size of the exposed population (545/1,291 = 43%), that the risk analysis be done by using the unexposed population as an internal reference group. The crude mortality ratio (CMR) for MM for the exposed population (51/291 = 0.04) is 0.4% (95% CI, 0.1-0.9), and CMR for the unexposed population (3/554=0.005) is 0.5% (95% CI, 0.1-1.6). The relative risk is 0.72 (95% CI, 0.17-2.98; Fisher two-tailed p-value = 0.70). When the analysis is limited to males as all the MM deaths were male, the relative risk is 0.62 (95% CI, 0.15-2.59, Fisher two-tail p-value = 0.45). The CMR analysis indicates that the MM risk in benzene-exposed workers is actually not greater than that of the unexposed. The marginally lower difference seen here is not significant. These results, like others reported in the literature, reinforce the im- portance of examining an internal reference group analysis when evaluating the po- tential risks associated with occupational chemical exposures. Analytic use of an in- ternal reference population reduces bias from the healthy worker effect or other risk factors associated with employment.

482 Expression of DNA Repair Genes in Breast Cancer Tumors from Puerto Rican Women.
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DNA repair is a critical defense system in the human body aimed at protecting the integrity and stability of the genome from the harmful effects of cancer-causing agents. Specific genetic alterations in DNA repair genes have been clinically associ- ated with risk, survival, aggressiveness, and treatment outcome of breast cancer (BC). Even though diminished DNA repair capacity is recognized as a risk factor for BC, no studies to date have identified gene expression profiles in tumors from Puerto Rican women. Study participants were women with histopathologically confirmed primary BC (n = 35) and normal breast tissue obtained during cosmetic surgery (n = 2). Tumors were snap-frozen in liquid nitrogen and RNA was extracted. Microarrays data were obtained from Affymetrix Plus 2.0 chip to assess whole genome gene expression. The assessment of differentially expressed genes was performed using Permuting method with 5% false discovery rate. Genome-wide gene expression levels were initially measured using the Affymetrix Plus 2.0 array, and DNA repair genes showing significant changes in expression levels were vali- dated by RT-PCR. Most of the DNA repair genes were over expressed. Twenty- three candidate DNA repair genes were found to be differentially expressed (in- cluding PARP-1, therapeutic target). Of these 21 were overexpressed and 2 were underexpressed (p<0.001). This is the first report of DNA repair candidate genes in Puerto Rican women with BC, a population with mixed ancestry. Supported by grants 506 GM000823-20, 1SCAI57250-2 and 5US56CA126379 from the NCI Center to Reduce Health Disparities and NIH-MBRS Program (NIGMS).

483 Investigation of Route-Dependent Exposure and Metabolism of Bisphenol A (BPA) in Neonatal Mice following Oral and Subcutaneous Administration of ‘H-BPA.
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Orally administered bisphenol A (BPA) undergoes efficient first-pass metabolism in the liver to produce inactive metabolites, including BPA-glucuronide (BPA-G) and BPA-sulfate (BPA-S). The current study was conducted to evaluate the pharmaco- kinetics of BPA and its conjugated metabolites, specifically BPA-G and BPA-S, in juvenile mice following a single oral or subcutaneous (SC) administration. This study had 3 phases: 1) Mass-balance phase in which effective dose delivery proce- dures for oral or SC administration of ‘H-BPA to postnatal day (PND) 3 mouse pups were developed; 2) Pharmacokinetic phase during which systemic exposure to total ‘H-BPA-derived radioactivity in female PND 3 mice was established; and 3) Metabolite profiling phase in which two groups of 50 female PND 3 mice received either a single oral or SC dose of ‘H-BPA at 400 μg/kg bw with = 4.4 mCi/kg bw of radioactivity. Blood was collected from 5 pups/route/time point at 5, 10, 20, and 30 minutes, and 1, 2, 3, 4, 6, and 24 hours post-dosing and processed to plasma. The plasma samples were pooled by group and time point and profiled by HPLC with fraction collection. Fractions were analyzed for total radioactivity and data used to generate radiochromatograms and integrate individual peaks. The identity of the BPA, BPA-G and BPA-S peaks was confirmed using authenticated standards and LC-MS/MS analysis. Metabolic profiles and key parameter as AUC0-24h and Cmax were derived for free-BPA, BPA-G and BPA-S. The result of this study re- vealed that female PND 3 mice have metabolic capacity to metabolize BPA to BPA- G, BPA-S and other metabolites after oral and subcutaneous administration and that sys- tematic exposure to free-BPA is route-dependent as the plasma concentrations were lower following oral administration compared to SC injection.

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Disposition and Kinetics of Tetrabromobisphenol a (TBBPA) in Female Wistar-Han Rats.


Tetrabromobisphenol A (TBBPA) is the brominated flame retardant with the largest production volume (~150,000 tons/year), representing nearly 60% of all worldwide demand for brominated flame retardants. TBBPA is used in printed circuit boards, ABS plastic casings, and laminates. Studies were conducted to characterize the disposition and toxicokinetic profile of TBBPA in female Wistar-Han rats following single oral bolus (25, 250 or 1,000 mg/kg) or intravenous (25 mg/kg) administration. All dosing solutions provided 50 μg/kg of [14C]-labeled TBBPA.

Following oral administration of [14C]-TBBPA the primary route of elimination of radioactivity in feces; dose recoveries in 72h were 95.7±3.5%, 94.3±3.6% and 98.8±2.2%, respectively. After a single IV administration of 25 mg/kg of [14C]-TBBPA, ~10% of the administered radioactivity was eliminated in the feces within 6h. Recoveries in urine ranged from 0.2-2.0% of dose. Less than 0.1% of the administered [14C]-radioactivity was detected in tissues collected at 72h following oral doses. Preliminary toxicokinetic profiles were estimated based on total [14C]-radioactivity detected in whole blood. Oral dosing (250 mg/kg) resulted in a rapid absorption of compound, with an apparent Cmax occurring at 3h post-dose and the subsequent distribution phase was consistent with a one-compartment model. Following IV administration (25 mg/kg), [14C]-radioactivity concentrations in whole blood decreased rapidly with less than 2% of the administered radioactivity remaining at 6h post dose. The time-concentration profile for total [14C]-radioactivity following IV administration of [14C]-TBBPA was consistent with a two compartmental model. These early results suggest low systemic bioavailability. These data indicate that TBBPA has a similar ADME/TK profile in female Wistar-Han rats.

These data indicate that TBBPA has a similar ADME/TK profile in female Wistar-Han rats. Toxicokinetics, NCI at NIEHS, Research Triangle Park, NC.

The model was calibrated using published BPA studies on in vitro hepatic and intestinal metabolism, in vivo biliary excretion of BPA metabolites, and in vivo serum time course data after oral and intravenous administration of BPA. Metabolism of BPA in the small intestine was predicted to substantially reduce the oral bioavailability of BPA and enterohepatic recirculation of BPA metabolites was predicted to prolong the systemic levels of BPA and its metabolites. The dosimetry of the aglycone BPA was age-dependent, in part, because of immature phase II metabolism in neonatal rats.

Human exposure to multiple pyrethroid insecticides may occur because of their wide use on crops and for residential pest control. To address the potential risk from exposure to pyrethroids, it is important to understand their toxicity and disposition in target organs such as the brain and surrogates such as the blood. The objective of this study was to compare motor activity with pyrethroid concentrations in blood and brain of rats after oral administration of a pyrethroid mixture. Male Long-Evans rats were dosed with a mixture of β-cyfluthrin (12.9% of dose), cypermethrin (28.8%), deltamethrin (3.4%), esfenvalerate (2.7%) and cis- (20.9%) and trans-permethrin (31.3%) in corn oil at one of seven doses (maximum total pyrethroid dose – 74.4 mg/kg). From 2 to 3 h post-administration of the treatment motor activity was assessed. At 3.5 h, blood and brain were collected and analyzed for parent pyrethroid using HPLC-tandem mass spectrometry. There was a linear dose-related increase in concentrations of the pyrethroids in blood and brain. Cypermethrin (56-62%) and cis-permethrin (57-70%) were the predominant pyrethroids detected in blood and brain, respectively, at all dose levels. The pyrethroids with the lowest percentage in tissue were trans-permethrin (0.6-2.7%) and β-cyfluthrin (0.4-1.2%) in blood and deltamethrin (1.1-3.5%) and esfenvalerate (2.4-4.7%) in brain. The approximate ED30 for decrease in motor activity was 300 ng/mL and 200 ng/g of total pyrethroid in blood and brain, respectively. More research is needed to understand the dosimetry of pyrethroids in blood and brain of rats and how this relates to neurotoxic effect. (This abstract does not represent U.S. EPA or NIEHS/NTP policy).

Investigation of BPA-Glucuronic Acid As a Substrate for Human BCRP, MRDR1, MRPR2 and MRPR3 Transporters.

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Bisphenol A (BPA) is a chemical used in plastic manufacturing that is present in polycarbonate bottles, food containers, and resins that line metal cans. Exposure to BPA occurs mainly through consumption of food products that are in direct contact with plastics and polycarbonates. Recently, BPA exposure in humans has been linked to the development of obesity and diabetes, as well as having effects on reproductive health and cancer. BPA is metabolized to BPA-glucuronide (BPA-gluc) and BPA-sulfate by Phase II enzymes and is eliminated by ATP-binding cassette (ABC) transporters. In humans, BPA-gluc is predominantly excreted through urine, whereas in rodents, it is excreted through bile into feces. One class of ABC transporters, multidrug resistance-associated proteins (MRPs) is thought to be involved in BPA clearance. There are nine human MRPs (MRP1-9), and mouse orthologs exist for all human MRP genes except MRP8. Multidrug resistance-associate protein (MRDR1) and breast cancer-resistant protein (BCRP) are also possible transporters for BPA metabolites. However, despite numerous studies, the mechanism of BPA metabolite clearance remains unclear. In this study we examined whether BPA-gluc is a substrate for human xenobiotic transporters BCRP, MRDR1, MRPR2, or MRPR3. Membranes expressing human BCRP, MRDR1, MRPR2, or MRPR3 were used in a colorimetric ATPase assay. The amount of inorganic phosphate (Pi) released from BPA-gluc-stimulated ATP hydrolysis was measured. Preliminary data suggest that BPA-gluc is a substrate for MRP3 (Km: 48.2 μM). Furthermore, BPA-glucuronide is a potential activator of MRP2 at low concentrations, with an ATPase transport activity >30% at 0.046 μM, but inhibits transport at higher concentrations. Low to negative percent ATPase activation values for MRDR1 strongly suggest that MRDR1 is inhibited by BPA-glucuronide. As differences exist between rodents and humans, identification of the ABC transporters involved in BPA clearance will be an important tool in the pharmacokinetic assessment of BPA.

Partition Coefficients of Deltamethrin (DLM) and Cis-Permethrin (CIS) in Male Sprague-Dawley Rats.

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Pyrethroid insecticides are widely used to control a wide variety of pests in and around homes, food handling establishments, in mosquito control, and in agriculture. Partition coefficients are essential to the construction of PBPK models, because of their role in determining systemic distribution. The main aim of this study was to determine tissue/blood partition coefficients (Ktp) of two commonly used pyrethroids, DLM and CIS. In vitro plasma to RBC partition coefficients, as determined by HPLC analysis, ranged from 0.7 to 1.0 for DLM, and 1.1 to 1.7 for CIS. The steady state levels of both DLM and CIS were obtained in vivo by constant infusion (0.36 mg/hr) using subcutaneous implantation of Alzet™ pumps, combined with oral loading doses of 34 mg/kg for DLM, or 150 mg/kg for CIS given 4 hr after the implantation of the pump in male rats. The time to reach steady-state was determined by analyzing blood samples collected by tail vein puncture 24, 48 and 72 hr post loading. Rats were sacrificed at 72 hr and blood and tissues collected and analyzed for DLM and CIS using a modified GC-MS method. DLM levels in plasma after 72 hr of constant infusion were 960 ng/mL, compared to 222 ng/mL in brain, 272 ng/mL in liver, 132 ng/mL in muscle and 1,384 ng/mL in fat. This corresponded to Ktp values of 0.23 for brain, 1.45 for fat, 0.28 for liver and 1.04 for muscle. In contrast, the CIS level in plasma was 78 ng/mL, compared to 43 ng/mL in brain, 27 ng/mL in liver, 107 ng/mL in muscle and 1,625 ng/mL in fat. These corresponded to Ktp values of 0.55 for brain, 20.78 for fat, 0.34 for liver and 1.37 for muscle. These data show that both pyrethroids are sequestered in fat, and that differences exist in Ktp for DLM and CIS, as Ktp values were generally higher for CIS. Supported by the Council For Advancement of Pyrethroid Human Risk Assessment.
Atrazine (ATR) is a widely used chlorotriazine herbicide that is commonly detected in the environment and in human specimens, including pregnant women’s urine, umbilical cord blood and breast milk. To help address concerns about reported adverse effects of ATR exposure during development, gestational and lactational PBPK models for ATR in the rat were developed. The models accounted for potential differences in the metabolism of ATR in dams, fetuses, and neonates and between single and repeated daily oral exposures and incorporated binding of ATR and its major metabolite deisopropylatrazine (DACT) with target tissue (maternal/neonatal brain and fetus), plasma proteins, and red blood cells. Model predictions correlate well with recently reported measured data on the concentrations of ATR and its metabolites in maternal/neonatal plasma, tissue, fetus, and milk following repeated daily oral exposure to the dam during gestation, lactation, or both, including with data from a study (lactational exposure) not used for model calibration. The model simulations indicate that: (1) the fetus is exposed to ATR and DACT at levels that are similar to maternal plasma levels, (2) the neonate is exposed mostly to DACT at levels about two-thirds of the maternal plasma DACT levels and (3) gestational carryover of DACT greatly affects neonatal dosimetry up until mid-lactation. Hence, excessive exposure to ATR and/or its metabolites during pregnancy or early lactation may be of particular concern. These models provide insights into designing and interpreting early life toxicology and pharmacokinetic studies with this herbicide and could be used in fetal and neonatal tissue dosimetry and for improvement of ATR’s exposure assessment.

Characterization of the Fate of β-Hexabromocyclododecanec (HBCD) in Mice.


1,2,5,6,9,10-Hexabromocyclododecane (HBCD) is a high production volume cycloaliphatic used primarily as an additive flame retardant in polystyrene foam building materials. Commercial HBCD mixtures contain three major stereoisomers, alpha (α), beta (β) and gamma (γ), at a typical ratio of 1.2:0.6:5.2. Toxicity (e.g. developmental neurotoxicity, immune effects, liver hypertrophy, and endocrine disruption) in rodents exposed to HBCD mixtures may be affected by differential kinetics of the isomers. Previous work from our laboratory demonstrated that α-HBCD has greater bioavailability and potential for accumulation in mice than γ-HBCD. The present investigation provides comparative data for β-HBCD to support toxicological evaluations of HBCD mixtures. In these studies, a single dose of [14C]-labeled β-HBCD (3 mg/kg), administered orally, was absorbed rapidly in the female C57BL/6 mouse. The Cmax for β-HBCD-derived radioactivity in blood and other assayed tissues, except adipose, was observed 3 hours following gavage. Approximately 8% of the total dose was eliminated in urine and feces by 24 hours. The extent of dose absorption was c. 85% based on HPLC analysis of feces extracts and comparison of oral and iv excretion data. β-HBCD-derived metabolites were excreted in urine and feces. Approximately 8% of the total dose was excreted in feces as γ-HBCD. Oral administration of either 30 or 100 mg/kg of β-HBCD resulted in initial slower changes in γ-HBCD concentration; however, cumulative excretion data were similar across the dosing range 4 days following gavage. 2-day cumulative excretion data in mice of these most were highest in adipose and liver. β-HBCD-derived radioactivity accumulated in these tissues following four consecutive daily doses of 3 mg/kg by gavage. In conclusion, β-HBCD, like γ-HBCD, was extensively metabolized, rapidly excreted, and had less potential for accumulation in tissues over time in female C57BL/6 mice than α-HBCD. (Supported in part by Intramural Research Program of NIH/NCI; abstract may not reflect official NIH policy.)

Perfluoralkyl sulfonates (PFASs) are a group of persistent environmental contaminants that have been detected in wildlife and human serum. Based on estimations from retired fluorochemical production workers, the elimination half-lives of certain PFASs, such as perfluorohexanesulfonate (PFHxS) and perfluorooctanesulfonate (PFOS), are very long (several years). Pharmacokinetic studies in animal models indicate that the long half-lives of these compounds are due to slow renal clearance and strong hepatic accumulation. In previous studies we have demonstrated certain organic anion transporters expressed in the kidney and the liver are involved in the disposition of another family of perfluoralkyl substances, perfluoralkyl carboxylates (e.g. PFOS) in rats. However, it is not known whether these transporters also substrates for the same drug transporters. Therefore, we wanted to test whether the organic anion transporters involved in the disposition of PFASs are also responsible for the disposition of PFOS. We used HEK293 cells expressing rat OAT1, OAT3 and OATP1A1 and measured the uptake of model substrates in the absence and presence of PFASs with different carbon chain lengths, namely perfluorobutanesulfonate (PFBS), PFHxS and PFOS. PFHxS showed the strongest inhibition for both rat OAT1 and OAT3, while rat OATP1A1 was only inhibited by PFOS. Direct uptake determination of these PFASs demonstrated that rat OAT1 and OAT3 can transport PFBS and PFHxS while rat OAT1 and OATP1A1 might transport PFOS. In conclusion, our results suggest that the same families of transporters for the rat that are involved in the disposition of PFASs are also involved in the disposition of PFBS, PFHxS and PFOS.
moM. 7:3 Acid was eliminated with calculated half-lives of 1-7 hrs in male and fe-male rats. The elimination of PFOA showed distinct gender differences with half-lives of 6-8 hrs in female rats and no measurable decline in male rats over the 18-h period of depuration. For both fluorotelomer alcohols, the major terminal acids formed metabolically in rats were 5:3 Acid and PFHxA from 6:2 FTOH and 7:3 Acid and PFOA from 8:2 FTOH. This study demonstrates rapid bioelimina-tion of inhaled fluorotelomer alcohols from plasma.

494 Absorption, Distribution, Metabolism, and Elimination of [14C] 6:2 Fluorotelomer Alcohol in the Rat.
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Fluorotelomer alcohols (FTOHs; F(CF2)6C2OH, x=6, 8, or 10) are used in the manufacture of specialty fluorinated surfactants and polymers. Fluorotelomer manu-facturers are moving away from raw materials that are potential precursors to per-fluorooctanoic acid (PFOA), such as the higher fluorotelomer alcohol homologues, to products based on 6:2 FTOH as a raw material. To better understand 6:2 FTOH biological fate, the absorption, distribution, metabolism, and elimination of [1,2-14C] 6:2 FTOH was investigated following a single oral dose administration of 5 and 125 mg/kg to male and female rats. The maximum concentration of total ra-dioactivity in plasma following oral dosing at 5 and 125 mg/kg occurred by 2 h post dose in both male and female rats. The plasma terminal elimination half-life values for total radioactivity were approximately 79 h in male rats at both dose lev-els and 78 and 63 h in female rats following a 5 and 125 mg/kg dose, respectively. Terminal elimination half-life values in red blood cells were approximately twice that in plasma, ranging from 121 h to 160 h. The internal dose as measured by area under the concentration-time curve to infinity (AUCinf) was similar for both male and female rats. The increase in AUCinf was slightly less than dose proportional from 5 mg/kg to 125 mg/kg. Preliminary material balance and tissue distribution data following an oral 5 mg/kg dose suggests that at 7 days postdose, less than 5% of the administered radioactivity was present in tissues with the highest concentra-tions occurring in fat, adrenals, and thyroid. The majority of the [1,2-14C] 6:2 FTOH dose was excreted in feces with 65% and 50% excreted in male and female rats, respectively. Renal excretion was also a significant elimination pathway with 10% and 18% excreted in urine in male and female rats, respectively.

495 Analysis of tris(2-Chloroisopropyl)phosphate Metabolites in Rat Plasma for Toxicology Studies.
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Tris(2-Chloroisopropyl)phosphate, TCP, is an organophosphate compound used as a flame retardant and plasticizer especially in polyurethane foam in furniture and in home insulation. Due to its environmental prevalence and resulting human ex-pposure, TCP is under study by the NTP. In support of NTP toxicological studies, MRIGlobal developed methods to analyze TCP and two major metabolites, Mono (MCPP, 2 isomers) and Bis 2-chloroisopropyl phosphate (BCPP, 3 isomers) in rodent plasma and blood. Because of its similarity to Tris(2-chloroethyl)phos-phate, BCPP was expected to be the major TCP metabolite. During the analytical method development, multiple isomers of MCPP were observed and hence were included in the method. The 5 isomers of the two metabolites were characterized using synthetic standards following derivatization with BSTFA/pyridine prior to analysis by GC/MS without using an internal standard. The ions monitored in the selected ion monitoring mode were for MCPP, m/z 211, 227, and for BCPP, m/z 155, 171, 197. The quantitation of MCPP and BCPP was achieved without derivatization using LC/MS/MS following extraction of 100 μL of plasma with 200 μL acetonitrile, re-moving acetonitrile by evaporation, and reconstituting the residue in 100 μL of 2% MeOH in water containing 50 mM tributylamine, TBA (pH 5 with formic acid) with internal standard dibenzylphosphate. The LC/MS analysis used a phenyl-hexyl column, which resulted in good linearity with a preliminary limit of detection value of 5 to 10 ng/mL of plasma ranging to over 500 ng/mL. Multiple MCPP and BCPP isomers were quantified as one MCPP and one BCPP peak. Plasma samples from rats and mice from single and multiple exposures to TCP were analyzed to show the applicability of this method. The data show the presence of MCPP and BCPP at levels significantly higher than the parent TCP.

496 In Vitro-In Vivo Extrapolation of 7-Ethoxycoumarin Metabolism Using 3D-Organotypic Liver Bioreactor.
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A liver bioreactor system was developed to simulate the metabolism and clearance of compounds similar to the liver in vivo to provide better prediction of in vivo clearance and metabolism profiles of compounds from in vitro data. Primary rat and cryopreserved human hepatocytes were maintained in a RealBio D475™ Culture System for approximately a month. General metabolic and liver-specific functions and cellular damage of rat/human hepatocytes inside liver bioreactor were moni-tored. Stable levels of biomarkers and related functional endpoints were achieved after a 7-10 day period for both rat and human liver bioreactors, respectively (glucose 64.6 – 86.6/152 – 182 mg/dL; lactate 0.6 – 1.9/0.1 – 3.7 mmol/L; pH 7.4 – 8.0/7.4 – 7.5; albumin 0.8 – 1.0/1.1 mg/mL; total protein 7.5 – 8.8/1.8 – 20 mg/mL; ALT 4.9 – 13.6/6.0 – 9.9 U/L; AST 6.8 – 25.4/18.9 – 53.1 U/L). The activities of both CYP2B2, CYP2C11 and CYP3A were monitored from the production of 16β-OH testosterone, 16α-OH testosterone and 6β-OH testosterone, respectively. Rat hepatocytes at 28 days in a liver bioreactor and treated with 5 μM dexametha-sone for 24hr showed 3-fold increase in CYP3A activity. Metabolism of 7-ethoxy-coumarin (7-EC) was characterized in the rat liver bioreactor. 7-Hydroxycoumarin was the predominant metabolite produced, which was further metabolized to the glucuronide and sulfate conjugates. The depletion of 7-EC showed mono-exponen-tial decay with an estimated intrinsic clearance of 113.3 μL/min/106 cells. Scaling of the in vitro C1int to in vivo C1int yielded in vivo prediction of 135:98 mL/min/rat body weight (250g). In conclusion, hepatocytes maintained in a 3D organotypic liver bioreactor showed stable liver functional capacity and intrinsic clearance over an extended period. This system may help resolve current limitations in assessing the clearance mechanisms and long-term effects of compounds and their major metabolites on chemical-induced liver toxicity.

497 Enhanced Intranasal Delivery of Gemcitabine to the Central Nervous System.
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Delivery of therapeutics to the brain to treat neurological diseases is a challenge due to impenetrable nature of the blood brain barrier (BBB). Intranasal (IN) drug ad-ministration is a non-invasive approach for rapid direct drug delivery from the nose to the central nervous system (CNS), thereby minimizing systemic exposure. The current study focuses on a strategy to enhance the delivery of the nucleoside drug gemcitabine (GEM) to the CNS via IN administration. Our approach takes advantage of the fact that the BBB and olfactory epithelial (OE) tight junctions (TJs) share many proteins in common. We hypothesized that by transiently increasing the permeability of nasal epithelial tight junctions using a BBB permeabilizer paverine (PV), we will increase the GEM crossing the brain extra-cellular fluid (BEFC) following IN delivery, with the goal of delivering therapeutic concentrations of nucleoside drugs to the CNS. Experimental methods included IN administration of fluorescein isothiocyanate-dextran beads (FD4), in-vitro GEM recovery, in vivo brain microdialysis for BEFC collection, HPLC analysis to measure GEM in BEFC, histopathology, and western blot analysis. Distribution studies with FD4 showed significant deposition in the ethmoid turbinates, suggest-ing drug uptake through OE. Clinically-relevant doses of PV (up to 1.4% IN) did not cause histological evidence of cytotoxicity or inflammation in nasal epithelia, lung, liver, spleen, or kidney. Pharmacokinetics of GEM in BEFC showed area under the curve (AUC) = 5.55±0.84 ng.h/ml for PV (1.4%) and GEM (50mg/kg) treated mice, compared to the AUC of 0.29 ng.h/ml for GEM without PV treatment. Western blot analysis suggested that IN PV treatment increased permeability through OE TJ by transiently decreasing the levels of TJ protein occludin. Thus, it appears that transient permeabilization of nasal epithelial TJs provides a non-inva-sive means to enhance delivery of nucleoside drugs to the CNS.

498 Altered Irinotecan Pharmacokinetics in Diet-Induced Obesity.

Purpose: Irinotecan (CPT-11) is a topoisomerase I inhibitor that has been shown to be highly effective in treatment of variety of cancers. It has recently been shown that CPT-11 administration is associated with liver toxicity and this effect is com-pounded by baseline obesity. It was found that patients with a BMI index of >25
were twice as much susceptible to developing liver toxicity than patients with BMI index of <25. CPT-11 metabolizes to SN-38, which then undergoes glucuronidation by uridine glucuronosyl transferase (UGT) 1A1 to form SN-38 glucuronide (SN-38G). Excess accumulation of the toxic metabolite SN-38 is known to cause fatal diarrhea in cancer patients. We hypothesize that accumulation of SN-38 is associated with increased liver toxicity of CPT-11 in obesity.

Methods: For metabolism studies, liver S9 fractions were prepared from diet-induced obese (DIO, 60% fat) and lean mice (10% fat). UGT1A-mediated metabolism of SN-38 was determined in liver S9 fractions. For pharmacokinetic studies, mice were injected with a single oral dose of 10 mg/kg CPT-11 and blood and feces samples were collected from 0-8 hr. Samples were analyzed for CPT-11 and SN-38 concentrations using LC-MS/MS. Liver tissues were harvested for real-time PCR studies. The mRNA and serum TNFγ levels were measured in liver and plasma samples, respectively.

Results: We found that the rate of formation of SN-38G was 2 fold lower in the DIO mice compared to the lean controls. This corresponded with reduced expression of UGT1A in these DIO mice. We also observed significant differences in transporter inhibition profiles may provide a selection criterion when both MRP2 and 4. Cyclosporine, vincristine and vinblastine only inhibited MRP2, M). With regard to the apical transporters, cy-

499 Interactions of Immunosuppressants with Human Organic Anion Transporters 1 and 3 and Multidrug Resistance Proteins 2 and 4.

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Renal proximal tubule transporters can play a key role in excretion, pharmacokinetic interactions and toxicity of immunosuppressant drugs. Basolateral organic anion transporters (OATs) and apical multidrug resistance proteins (MRPs) contribute to active tubular uptake and urinary efflux, respectively. Combining immunosuppressants during therapy may lead to drug-drug interactions occurring at the transporter level, resulting in toxicity. We studied the effect of different immunosuppressants on OAT1- and OAT3-mediated uptake of [3H]-methotrexate ([3H]MTX) in cells, and on ATP-dependent [3H]MTX transport in membrane vesicles isolated from HEK293 cells over-expressing human MRP2 and MRP4. For the uptake transporters, we found that cyclosporine, dexamethasone, azathioprine and 6-mercaptopurine did not affect either transporter. Cytarabine, vinblastine, vincristine, hydorchloroic acid and mitoxantrone significantly inhibited OAT1 (p<0.05 at 10 μM), whereas these compounds did not inhibit OAT3. In contrast to other compounds, mycophenolic acid (10 μM) inhibited OAT3 more effectively than OAT1, reducing [3H]MTX uptake by 86 ± 4% and 52 ± 5% of control, for OAT3 and OAT1 respectively. Subsequent studies showed that the IC50 of mycophenolic acid for OAT3 was 4.3 μM (95% CI: 2.7-6.9 μM), which is close to its reported Cmax, unbound (~1 μM). With regard to the apical transporters, cytoplasmic, hydorchloroic, tacrolimus and mycophenolic acid inhibited both MRP2 and 4. Cyclosporine, vincristine and vinblastine only inhibited MRP2, while 6-mercaptopurine only inhibited MRP4. In conclusion, immunosuppressants may alter renal transport activity. From a drug safety perspective, the observed differences in transporter inhibition profiles may provide a selection criterion when combining immunosuppressive drugs.

500 Accumulation of β-α-Methylamino-L-alanine in Tissues Following Repeat Oral Administration to Harlan Sprague-Dawley Rats.

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β-α-Methylamino-L-alanine (L-BMAA) was nominated to the National Toxicology Program for toxicological assessment based on widespread environmental distribution and evidence that the potentially neurotoxic compound may accumulate in CNS tissue. Data describing metabolism and disposition of L-BMAA are needed to support placement of NTP toxicity study. Male Harlan Sprague-Dawley rats were dosed by gavage with [1,2-14C]-L-BMAA (1 mg/kg/d) for 1, 5, and 10 consecutive days. Excreta and tissues were collected for up to 72 hours after the final dose. The majority of 14C was recovered as 14CO2 (50-60%) across 10 days of dosing and was eliminated in urine. HPLC peaks of 14C over time showed multiple polar metabolites and L-BMAA was not detected. Over 10 days of dosing 14C continued to accumulate in tissues including the brain. After single and up to 10 day repeat doses the majority of the 14C recovered in tissues was found in liver, adipose, muscle, and skin; <0.01% dose was recovered in the brain. Accumulation rate of brain 14C over 10 days exceeded elimination by 3 fold. On days 1,5 and 10 a majority of 14C in brain tissue 24 h following the final dose was recovered in the protein after exhaustive extraction. The nature of BMAA associated with brain protein was investigated in vitro. Incubation of [1,2-14C]-L-BMAA in vitro with rat brain homogenates released 14CO2 and 14C was incorporated into protein. HPLC analysis of in vitro protein hydrolysates showed that [14C]-L-BMAA itself and other 14C-equivalents made up the radioactive components. Thus accumulation of L-BMAA and its equivalents in brain tissue may be due in part to incorporation into protein. Brain tissue from repeat dose administration in rats is currently being analyzed to investigate the nature of the radiolabel associated with the protein. This work was conducted for the NTP under NIHES Contract N01-ES-75562.

501 Comparative Nonclinical Ocular Tissue Distribution of the Visual Cycle Modulator (VCM) Emixustat in Rats, Beagle Dogs, and Cynomolgus Monkeys.

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Emixustat HCl is a novel, non-retinoid VCM that inhibits a key step in regeneration of 11-cis-retinal by retinyl ester isomer RP665. Emixustat is currently in late Phase 2 clinical development as an orally administered treatment for geographic atrophy associated with dry age-related macular degeneration (AMD). In ADME studies, very low plasma concentrations of emixustat are seen as it is extensively metabolized, producing pharmacologically inactive metabolites. Interestingly, the pharmacological activity of emixustat has been shown to be prolonged beyond the time at which its plasma concentrations are no longer measurable. Ocular distribution studies using 14C emixustat were undertaken in rat, dog, and monkey to better understand the time course of emixustat ocular tissue exposure in relation to observed pharmacology activity in animal models. Pigmented and albino rats received single oral doses of 1 mg/kg, and were prepared for QWBA, or a repeated oral dose (QD for 7 days) and were prepared for ocular dissection. Dogs received 0.3 mg/kg as a single or a repeated oral dose (QD for 7 days). Monkeys received 0.9 mg/kg/day as a single or a repeated oral dose (TID for 7 days). Eye levels were observed for up to six weeks postdose. The mean recovery of radioactivity was >95% in rat and dog, and >90% in monkey. The half life of removal of radioactivity from eye tissues was longer in all species than that observed in plasma. Consistent among the three species, emixustat parent molecule was the major component found in eye tissues. In all three species, the retina Cmax of emixustat were >50 fold higher than respective plasma Cmax on a ng-equiv/g basis. Although high levels of the major metabolites were observed in plasma, they were not observed in eye tissues. Preclinical ocular distribution results indicated emixustat parent is the predominant molecule in the eye responsible for pharmacological activity.

502 Development of Sustained Release Buprenorphine for Use As an Improved Analgesic in Toxicology Studies: Assessment of Formulation Pharmacokinetics.

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Acute toxicity safety testing procedures can involve animal pain and distress. U.S. regulatory agencies and the OECD recently adopted and updated procedures that incorporate the routine use of systemic anesthetics to avoid or reduce pain and distress for eye irritation testing procedures. Buprenorphine is recommended as a useful analgesic for such toxicity studies. However, buprenorphine requires a minimum of twice-daily dosing at 12 hour intervals to maintain effective analgesia. A study was therefore conducted to evaluate sustained release formulations to determine their usefulness for once-per-day or less frequent dosing. The pharmacokinetics of two sustained release formulations of buprenorphine were compared. Acute toxicity study results showed that buprenorphine was converted to metabolites in saline using male 2.2-2.5 kg New Zealand White rabbits. Sustained release formulations were prepared using N-methyl-pyrrolidone (NMP) or Triacetin as the vehicle. Following subcutaneous dosing, blood samples were collected at intervals to four days; plasma buprenorphine was determined using liquid chromatography with mass spectrometry. Both the NMP and Triacetin formulations produced higher plasma buprenorphine concentrations than the saline formulation at time points from 12 hours on. The Triacetin formulation also resulted in higher concentrations at earlier time points, as well as concentrations near or
above 0.1 ng/mL—a concentration previously associated with analgesic activity in other species—by breath, post-dosing. Both urine and breath concentrations were recorded, and did not show adverse effects of treatment with sustained release formulations. No abnormal clinical signs or local lesions were observed. These results suggest that the Tricaten sustained release formulation of buprenorphine can significantly reduce the dosing interval required and can be a useful replacement for twice-daily treatments.

503 Acute Rat Inhalation Pharmacokinetic Study of 10.9 kD Protein with or without Pegylation.

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Biologics are an evolving protein-based therapeutic approach in the pharmaceutical industry. As part of assessing hazards, an Occupational Exposure Limit (OEL) is developed internally for proprietary biologics to support worker safety. Biologics are considered to be less hazardous than small molecules due to suspected lower systemic bioavailability (BA) from the lung and degradation in the GI tract. To investigate fate after inhalation (Inh), SD rats were exposed to RGE (10.9 kD protein), RGE-PEG (40 kD PEGr10.9 kD protein) or saline by nose-only Inh for 1 hr. The pharmacokinetic (PK) profile was investigated in blood and bronchial alveolar lavage fluid (BAL). The measured experimental exposures were 42 and 48 mg/m3, corresponding to a calculated dose of 1.8 and 2.1 mg/kg, and a mean mass aerodynamic diameter of 1.8 and 1.9 μm for RGE and RGE-PEG, respectively. The systemic PK profile for RGE and RGE-PEG included AUC(INf) of 169 and 357 ng·h/mL; Tmax of 2 and 9 hrs; and t½ of 5.5 and 21 hrs, respectively. When compared to an IV study, systemic BA was 1.4 and 0.05%, respectively. From BAL levels, the left lung lobe contained 4.2 and 3.2% of the calculated dose at end of exposure and the lung t½ was 4.5 and 7.2 hrs, respectively. These results indicate that after Inh exposure, systemic BA was low, with the PEG-RGE being markedly less than the unmodified RGE. However, the overall AUC(INF) for both proteins was similar. Compared to systemic results, the lung PK for the two proteins were similar with respect to t½ and reduced percent of dose delivered to the lung. These PK results suggest that compared to small molecule drugs, there may be a decreased hazard for proteins, which would be demonstrated by applying these data-derived PK factors to the OEL calculation. Further studies on additional biologics would be needed to generalize across proteins.

504 Simulation of Perfluoroacid Accumulation in Humans Exposed to Perfluorotelomer Alcohol Products.


Perfluorochemical grease-proofing agents derived from perfluorotelomer alcohols are commonly used in fast-food packaging, and have been of concern because of the surfactant perfluorooctanoic acid (PFOA) and related perfluoroalkyl carboxylic acids (PFCas). S-glutathionylated, studies in the rat indicated that small amounts of PFCAs are also absorbed. These new toxicokinetic data and improved PBPK model for ethanol will facilitate the refinement of risk assessment for chronic inhalation exposure to low levels of ethanol. (Project funded by ANSES, France).

505 Uptake and Disposition in Humans of Hydrofluorocarbons (HFCs) Used As Refrigerants.

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A variety of hydrofluorocarbons (HFCs) have replaced the ozone-depleting chlorofluorocarbons (CFC) and hydrochlorofluorocarbons (HCFC) during the last decades. There are few data on uptake and disposition of HFCs in humans. In a series of experiments, we therefore exposed healthy volunteers to vapors of four commonly used HFCs, namely difluoromethane (HFC152a; 0, 200 and 1000 ppm), trifluoroethane (HFC143a; 500 ppm), tetrafluoroethane (HFC134a; 500 ppm) and pentafluoro propane (HFC245fa; 0, 100 and 300 ppm). In parallel, blood, salivary and urine samples were collected until next day and analyzed for the parent substance by head-space gas chromatography. Fluoride and other potential metabolites were analyzed in urine using an ion selective electrode and 19F-NMR analysis, respectively. All HFCs had similar toxicokinetic profiles in blood with a rapid initial increase of HFC and an apparent steady-state reached within a few minutes. The area under the concentration-time curves (AUC) of HFC152a and HFC245fa in blood was proportional to the exposure level, suggesting first-order kinetics. For all four HFCs, the inhalation uptake was low (less than 4%) and only minor amounts were excreted unchanged in breath and urine after exposure. No signs of metabolism were detected except a slightly increased urinary excretion of fluoride after exposure to 1000 ppm HFC152a. No other urinary metabolites were detected. The observed time courses in blood and breath could be well described with a physiologically-based pharmacokinetic (PBPK) model. In conclusion, the uptake and disposition of the four HFCs were consistent with the PCs determined in vitro and with zero or insignificant biotransformation.

506 Ethanol Toxicokinetics Resulting from Inhalation Exposure in Human Volunteers and Toxicokinetic Modeling.

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There is a lack of information and increased interest on the risks associated with chronic inhalation exposures to low levels of ethanol (< 1000 ppm). A physiologically based pharmacokinetic model (PBPK) for inhaled ethanol was previously developed based on exposed volunteers but only for levels above 5000 ppm. Uncertainty still remains about the validity of this model to predict the blood levels of ethanol (BE) for lower level exposures. This project aims to determine the BE resulting from exposure to low concentrations (<1000 ppm) in order to validate the PBPK model. Ten volunteers (5 men and 5 women) were exposed for 4 hr to vapors of ethanol (125, 250, 500, 750 and 1000 ppm) in resting conditions in an inhalation chamber. An additional exposure to 750 ppm that included 4 periods of 12 minutes of exercise at 50 W was performed. Blood samples and alveolar air were collected during and after the exposure. Results show that there is a linear relationship between the ethanol inhaled air concentrations and (i) BE (women: r² = 0.98/men: r² = 0.99), as well as (ii) ethanol concentrations in the alveolar air at end of exposure period (men: r² = 0.99/women: r² = 0.99). Furthermore, exercise resulted in a significant increase (2 to 3 times) in BE after each period of 12 min of exercise. Overall the model predictions were overestimated for very low levels (< 216 ppm for men and < 2300 ppm for women. At lower exposure concentrations, limiting the clearance to the liver compartment was insufficient to account for total ethanol clearance. Adjusting the model by adding extra-hepatic biotransformation of high affinity and low capacity associated with the richly perfused tissues allowed this model to fit adequately the low and high exposure level toxicokinetic data. PBPK model for ethanol will facilitate the refinement of risk assessment for chronic inhalation exposure to low levels of ethanol. (Project funded by ANSES, France).

507 Physiologically-Based Pharmacokinetic (PBPK) Modeling: Extrapolation from In Vitro to In Vivo, a Case Study Using a Novel Dermal Compartment.


PBPK models are useful extrapolation tools requiring time course data for calibration. A unique data set for rodents and humans was used with an existing PBPK model for octanol. The model predictions of the same compounds were extrapolated in vivo from in vitro extrapolation, an emerging application technique for PBPK modeling. By adding a dermal compartment, a refined rodent PBPK model for orally administered lindane was developed and optimized using time-course tissue concentration data for Wistar rats. This refined PBPK model also provided blood and skin partition coefficients (Phlodd = 1.72 and PskIn = 26.2); other partition coefficients were obtained from the literature. Next, a human in vivo model was extrapolated from the rodent model using physiological values from the literature. In addition, a novel in vitro model of human skin containing a follicular compartment was developed to...
improve the fit to the available in vitro absorption data. In vitro dermal permeability coefficient was estimated using two different methods: (1) a permeability coefficient of 0.013 cm/hr derived with the Potts-Guy equation; simulations generated with this coefficient matched the observed dermal permeability results in vivo; and (2) a permeability coefficient of 0.0043 cm/hr derived from the dermal in vitro model; simulations generated with this coefficient matched the observed dermal permeability results in vivo. The permeability value obtained using the in vivo PBPK model was 0.0060 cm/hr, which provided the best fit to the data. This result is between the in vitro and the Potts-Guy estimates, and PBPK simulations generated with this in vivo permeability fit through the averaged data points. In summary, PBPK modeling was found to be a powerful in vitro to in vivo extrapolation tool, particularly when time course datasets are available. (This abstract does not reflect EPA policy.)

508 Evaluation of Toxicity Adjustment Factors Used for the Risk Assessment of Chlorpyrifos Oxon in Drinking Water.

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In the Preliminary Human Health Risk Assessment for chlorpyrifos (CPF), EPA has derived toxicity adjustment factors (TAF) for chlorpyrifos oxon (Oxon), for both acute and chronic dietary scenarios, and 12 and 18, respectively, based on relative inhibition of RBC acetylcholinesterase (AChE) by Oxon, relative to CPF. However, a comprehensive evaluation of the biological effects of Oxon, following oral exposure, has shown that this test material undergoes complete first-pass metabolism, via chemical and/or enzymatic processes in the GI tract, GI tissue, portal of entry (i.e., do not directly affect other tissues). Oxon was not considered to have long-term clinical consequences, and would not be expected given the predicted, low level exposures to Oxon in drinking water. As a result of these analyses, it is proposed that the potency of Oxon be based on a relevant toxicity endpoint (e.g., brain effects seen at > 10 mg/kg) and not on RBC ChE inhibition, which is an indicator of low levels of Oxon that are not systemically available.

509 Effects of Leucine Administration on Plasma and Brain Levels of Other Amino Acids.


Leucine (leu), one of the essential branch chained amino acids (AAAs), has been shown to activate mTOR to increase protein synthesis, and for the brain, may provide enhanced cognitive memory deposition. However, little is known regarding leu kinetics. Investigations into the leu kinetics revealed that it was eliminated very fast from the blood after dosing, and leu levels in brain were higher than blood, indicating active transport of leu across the blood brain barrier (BBB). The objective of this work was to investigate the effects of leu administration on blood and brain levels of 20 AAAs, and the BBB role (if any) to leu kinetics. A rodent model was dosed leu via iv (5 & 12.9 mg/kg) for dose-response and time-course studies, and orally (319 mg/kg) for time-course analysis. Tissue leu levels were measured at various time points (5m– 6h, iv and 30m–4h, orally). Interestingly, more than 65% of all AAAs were increased in brain. At the 5 mg/kg dose, valine, lysine and proline in brain were increased, while threonine and methionine decreased (all relative to control, p<0.05). With oral dosing, 16 AAAs in brain were increased at the 1h time point. Isoleucine (time point 1 & 4h), lysine (0.5 & 1h), histidine (0.5 & 1h) and proline (all time points: 0.5, 1, 2, and 4h) in brain were increased significantly from control at p<0.05 or less. Brains to plasma AA ratios were increased following dosing for lysine, methionine and valine at 5 mg/kg, proline at 12.9 mg/kg and isoleucine and proline for oral dosing groups. Our results indicate that leucine administration initiate complex time- and dose-dependent responses in both plasma and brain levels of the other AAAs. These preliminary data show that neither the specific BBB transport system for AAAs nor the general BBB transport system for small neutral AA transporters for crossing the BBB, nor obvious general chemical or biological AA properties determine the brain levels and uptake behavior observed in these studies.

510 Screening for Substrates of the P-Glycoprotein Transporter.


P-glycoprotein (Pgp) is an ATP-dependent efflux transporter of xenobiotic compounds with broad substrate specificity. Pgp is an important component of the blood-brain barrier and frequently mediates resistance of tumors to chemotherapeutic agents. Attempts to predict the in vivo toxicity of environmental chemicals based upon in vitro screening test results can be erroneous if transport effects are not considered. In this study, a multi-tiered in silico/in vitro testing approach was used to predict Pgp substrates within a set of ToxCast chemicals. An in silico analysis using a support vector machine predicted 45% of the 280 chemicals to be Pgp substrates; these predictions matched the results of a separate in silico docking analysis. Only 5% of the chemicals with MW < 400 (and none with MW > 300) were predicted as substrates, while 87% with MW > 400 were predicted substrates. In vitro cell-based assays were used to further screen for Pgp substrates among chemicals with MW > 300. Dye efflux experiments were conducted with NIH 3T3 MDR1 cells that stably express human Pgp. A number of potential substrates were identified by this assay (including ivermectin and abamectin), as shown by their capability to interfere with efflux of Hoechst 33342 dye from the cells. A more definitive assay was used to test chemicals with known cytoxicity by comparing toxicity in the MDR1 cells and wild type 3T3 cells. Pgp substrates (i.e., chemicals that were less toxic to MDR1 cells) included captan, naled, pyriproxifen, thiophanate, thiocarb, abamectin, niclosamide, and rotenone. These results demonstrate that a systematic approach involving a combination of in silico and cell-based in vitro assays can successfully predict Pgp substrates within large chemical test sets. In vitro toxicants that are identified as Pgp substrates will require additional toxicokinetic assessment to correctly predict their potential in vivo toxicity. (This abstract does not reflect EPA policy.)

511 Transport Mediated Mechanism for Nucleoside Penetration of the Blood-Testis Barrier.

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The blood-testis barrier (BTB) is formed by tight junctions between Sertoli cells and prevents the entry of many therapeutics into the lumen of the seminiferous tubules (STs) shielding developing germ cells from chemical exposure. One drug class of HIV therapeutics, nucleoside reverse transcriptase inhibitors (NRTIs), can penetrate the BTB and be detected in human seminal plasma at concentrations higher than that of blood plasma. The purpose of this study is to determine the mechanisms by which NRTI drugs are transported across the BTB. Transport studies in isolated rodent seminiferous tubules using H3uridine (Kt 89.72uM) as a model nucleoside substrate indicate that seminiferous tubules take nucleosides almost exclusively via equilibrative nucleoside transporter 1 (ENT1). The IC50 for NBMPR, an ENT inhibitor, is 12.9 nM. Trans-epithelial transport of uridine by primary cultures of Sertoli cells can also be blocked by ENT1 inhibition. Blocking ENT1 function on the basolateral membrane also prevents uridine uptake into cells. These data correspond with immunohistochemical staining of rat testes showing ENT1 on the basolateral membrane, whereas ENT2 is on the apical membrane of Sertoli cells. This localization suggests that ENT1 acts as an uptake transporter and ENT2 may facilitate the efflux of nucleosides and NRTI drugs into the lumen of STs. Uridine transport can also be inhibited by NRTI drugs. We also demonstrate transepithelial transport of radioabeled NRTIs zidovudine (AZT) and didanosine (ddI) through primary Sertoli cells is partially blocked by ENT1 inhibition (88% and 67% for AZT and ddI respectively). These data indicate a novel ENT dominant mechanism for the transepithelial transport of nucleosides and NRTI drugs across the BTB.

512 Pharmacokinetics and Bioavailability Testing of Levofloxacin in Rats.

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Drug exposure of the antibiotic levofloxacin was examined in male and female Sprague Dawley rats following oral and intravenous administration. Pharmacokinetic modeling and determination of bioavailability were conducted following single dose treatment and subsequent measurements of levofloxacin levels in plasma. Dose levels studied were 50 mg/kg oral (PO) and 25 mg/kg intravenous.
There were no abnormal clinical signs observed following dosing, which might have otherwise affected the pharmacokinetic modeling. The systemic bioavailability of levofloxacin in rats after PO administration ranges between 33.7 - 42.7%, and appears to be lower in males. The volume of distribution (Vd) appeared to be higher in female than in male rats following IV administration. Other pharmacokinetic parameters appeared to be similar in both male and female rats for both routes of administration. Levofloxacin appears to distribute widely in rat tissues and possibly exhibits a large affinity for tissue proteins. The half-life of the drug is short (4.33 hr and 4.96 hr in males and females, respectively, following PO administration; 1.38 hr and 1.55 hr in males and females, respectively, following IV administration), probably due to a high systemic clearance. Levofloxacin single IV (25 mg/kg) and PO (50 mg/kg) doses were well tolerated by male and female rats. Mean (±SD) AUC0-inf values following IV administration were 11339.0 ± 572.3 hr·ng/mL and 9422.7 ± 1169.4 hr·ng/mL, in males and females, respectively, and following PO administration were 8051.4 hr·ng/mL and 10211.6 hr·ng/mL in males and females, respectively. Based on mean Tmax and terminal elimination half-life values, levofloxacin is rapidly absorbed and eliminated.

513 Comparative Disposition and Metabolism of 2, 2'-Dithiobisbenzanilide following Dermal, Oral, and Intravenous Administration to Harlan Sprague-Dawley Rats and B6C3F1/N Mice.

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2, 2'-Dithiobisbenzanilide (DTBBA) is used as a peptizing agent in tires and rubber products and occupational exposure to DTBBA may occur mainly through dermal contact. Potential systemic exposure via the dermal route was investigated in rodents and compared to intravenous and oral routes. A single dermal doses of [14C]DTBBA (4 mg/kg, in acetone) were applied (protected from oral grooming) to male Harlan Sprague-Dawley rats and B6C3F1/N mice and its disposition and metabolism 72 h after was compared to 4 mg/kg intravenous and gavage doses. Following a dermal dose of [14C]DTBBA in rats and mice, 10.9 ± 2.9 and 10.6 ± 2.9% of the dose was absorbed, respectively. Excretion of absorbed dose was via urine (18.4 ± 3.3, 34.7 ± 7.7 and feces (32.6 ± 10.7% and 49.5 ± 10.8%) in rats and mice, respectively. Following gavage administration in both species, the absorption was complete. Excretion of [14C] after rat intravenous doses was mostly via urine (73.1 ± 17.8%). After mouse intravenous, mouse or rat dermal or gavage doses -30% -40% was recovered in urine and feces. The percent dose in tissues after dermal dosing was 4.58 ± 1.92% for the rat and 0.98 ± 1.13% for mouse and in both species non dose site skin had the highest [14C] levels. Radioactivity remaining in tissues followed the trend of dermal > intravenous > gavage in rat and intravenous > gavage > dermal in mouse. DTBBA was not detected in urine. Urine metabolite profiles were qualitatively similar between species but the levels of some metabolites varied between dose routes. The structure of the predominant urinary metabolite was identified as thiobisbenzanilide-S-glucuronide by LC/MS/MS and 1H and 13C-NMR. This work was conducted for the NTP under NIEHS Contract N01-ES-75562.

515 Lipophilicity and Membrane Affinity of Munitions Constituents Using BioLipid Beads.

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Bioaccumulation of organic molecules into fat can be predicted by chemical lipophilicity which provides the highest systemic absorption of HMB and there is a difference in absorption between rats and mice. This work was conducted for the NTP under NIEHS Contract N01-ES-75562.
lead to i) a significant reduction in animal use (e.g., 32 instead of 48 rats in a 14-day dose-range finding study), ii) the abandonment of anesthesia for toxicokinetic blood sampling, and iii) better data as effect and exposure are gathered in the same animal leading to better assessments of drug-related effects.

The benefits of QTOF bioanalytical quantification for large therapeutic peptides (minimum drug optimization; generic method and non-targeted quantification) were confirmed and it can be used for TK & PK studies.

### 517 Using PBPK Modeling to Address Diurnal Variation and Age Differences in Hexavalent Chromium Toxicokinetics in Humans.

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A physiologically based pharmacokinetic (PBPK) model has been developed to describe the toxicokinetics of hexavalent chromium (Cr(VI)) in mice, rats, and humans. The PBPK model was used to support a human health risk assessment (HHRA) based upon mouse small intestinal (SI) tumors. Key factors contributing to the delivery of Cr(VI) to the SI were identified by sensitivity analyses, and include gastric pH, gastric transit time, and gastric reducing equivalents. These factors affect the rate of Cr(VI) reduction in gastric contents, with a higher delivery of Cr(VI) resulting from higher pH values (which causes slower rates of Cr(VI) reduction), shorter stomach transit times, and lower reducing equivalent concentrations. The PBPK model was used to address 4 important sources of variation. First, the model was used to account for normal diurnal variation in gastric lumen factors (e.g., in normal individuals, baseline gastric pH is typically between 1-3 between meals, but rises rapidly to levels of 5-7 at the start of a meal, then returning to baseline levels within 2-3 hours). Second, the model was used to simulate exposures to different age groups, including infants, children, youths, adults, and elderly, since some age groups (e.g., infants) normally exhibit higher pH values than adults. Age-specific differences in the key factors were incorporated in the modeling to estimate human equivalent lifetime average daily dose corresponding to point of departures determined for mouse SI lesions. Third, the model was used to assess risk to specific populations (proton-pump inhibitor users) that may be sensitive to Cr(VI) due to alterations in gastric pH. Lastly, the model was used to assess the impact of the timing of Cr(VI) exposure events on delivery of Cr(VI) to the SI with respect to their occurrence during or between meals. The implications to HHRA are quantified and discussed.

### 518 Advantages of Using Quadrupole Time-Of-Flight (QTOF) in the Bioanalysis of Large Therapeutic Peptides.


**Purpose.** MUC5AC13, hectarin, and calcitonin were selected as model compounds to demonstrate the advantages of QTOF in large therapeutic peptides bioanalysis for toxicokinetic (TK) and pharmacokinetic (PK) studies.

**Methods.** Samples were prepared by spiking MUC5AC13, hepticin or calcitonin in plasma. They were extracted using protein precipitation followed by solid phase extraction. UPLC was used for chromatography. MUC5AC13 and Calcitonin were analyzed on TripleTOF5600 with DuoSpray. A KinetexC18 column and 0.6ml/min flow rate in gradient conditions were used. For heptacin, electrospay Chip emitters TriVersaNanoMate on TripleTOF5600 with AtlantisT3 column at 0.5UL/min flow rate in gradient conditions were used.

**Results.** MUC5AC13 showed multiple charge states +2 and +3 as the most abundant ions in QTOF full scan. Summing +2 and +3 charge states with a 10Da extraction window was used for quantification, over 3.5 orders of linear range (0.4-2000ng/mL), a precision of <9.3% and an accuracy between 86-113% were obtained. In hepticin quantification, full scan and parent ion quantification showed interferences even with narrow extraction window (2-5mDa). The interference peaks might be in source fragments and thus eliminated by the first quadrupole (Q1). Hence, quantification of hepticin was performed in QTOF tandem mass spectrometry and summing +4 charge state isotomers of parent ion at a 10mDa extraction window and it showed 3 orders of linear dynamic range (2-2048ng/mL) a precision of <9% and an accuracy between 80-112%. Calcitonin quantification showed +3, +4 and +5 charge states as the most abundant ions. QTOF full scan and summing +3, +4 and +5 charge states of parent ion at a 10mDa extraction window were used for quantification. The observed LLOQ was 50pg/mL with 4 orders of linear dynamic range. The precision was better than 15% and the accuracy between 83-118%.

### 519 A New Strategy to Effectively Reduce Matrix Effect Caused by Phospholipids in Plasma Samples under Hydrophilic Interaction Liquid Chromatography (HILIC).


**Purpose.** Significant reduction of matrix effect due to phospholipids (PL) in plasma sample by using aprotic solvent to improve the performance of bioanalytical methods used for drug quantification in toxicokinetic and pharmacokinetic studies.

**Methods.** Rivastigmine was extracted using liquid-liquid extraction and injected using HILIC column with ammonium acetate / Acetonitrile (ACN) with a flow rate=0.8mL/min. Analysis was performed on ABSciexAPI31000 with electrospray(+). Lower limit of quantification=10pg/mL. PL were monitored at m/z 184. The PL extractability was evaluated by samples evaporated in polypropylene (PP) instead of glass tubes and reconstituted with ACN.

**Results.** Extracted plasma samples were reconstituted in mobile phase (MP blank) or in ACN blank and injected with post-column infusion of rivastigmine. The post-column infusion of MP and ACN blanks showed no suppression at the drug retention time. However, the baseline from the post-column infusion of MP blank dropped by 50% after seven injections. However, ion-suppression was not observed for the ACN blank. These results suggest that the late eluting suppressors are not reconstituted in ACN. PL peak was observed after seven MP blank injections suggesting that the late eluting suppressors in the MP blank are phospholipids. This peak was not observed in ACN blank. These results suggest that PL are reconstituted in mobile phase and elute as late eluting suppressors under the chromatographic conditions. However, PL are not reconstituted using ACN. The extracted blank plasma samples were reconstituted in PP tubes with ACN and compared to MP and ACN blanks reconstituted in glass tubes to determine if the PL are insoluble in ACN or if adsorption occurred on the glass tube surface. The PP ACN & MP blanks chromatograms were identical thus demonstrating that PL are soluble in ACN but were adsorbed on silica surface.

**Conclusion.** The reconstitution of extracted samples in glass test tube using 100% ACN successively eliminates the matrix effect caused by phospholipids.

### 520 Impact of Organic Solvent Additive on the Integrity of Plasma Samples over Time in Bioanalysis by LC-MS/MS.


**Purpose.** Evaluation of plasma integrity over time due to addition of organic solvent in quality control samples (QCs) for morphine-6-glucuronide (M6G) bioanalytical method that was validated to be used in pharmacokinetic studies.

**Methods.** The samples were extracted using solid phase cartridges. The injection of the samples was performed on XBridge Phenyl column (2.1x50mm, 5µ) in gradient conditions on Agilent1100 coupled with ABSciexAPI5000 in electrospray(+). The QCs were prepared with M6G alone and M6G + naltrexone (Co-Administered Drug - CAD) at 1.0% organic content vs. 1.5% respectively. M6G and M6G-D3 (IS) were monitored at m/z 462/201 and 465/289 respectively. Full scans and post-columns infusion were performed to evaluate the presence and determine the impact of interfering compounds.

**Results.** A significant decrease in signal (∼35%) was observed for M6G/IS over time for the QC containing naltrexone. A post-column infusion profile of these QCs showed the presence of suppressors which co-eluted with M6G/IS. Chromatographic separation was achieved between naltrexone and M6G/IS thus eliminating any possible suppression from naltrexone. Full scan experiments enabled the detection of two specific masses co-eluting with M6G/IS only in the QCs stored over time and containing naltrexone. QCs containing only the analyte and others containing the analyte with naltrexone were prepared in different plasma matrices. Some blank matrices were tested alongside with blank matrix containing different organic solvent percentage. The freshly prepared QCs did not show ion suppression; either in the presence of naltrexone or of the extra organic solvent. The long-term evaluation of the matrices in the presence of organic solvent (1.5%), showed two unknown compounds to appear over time and elute at the retention time of M6G thus creating a suppression zone.
Conclusion.
The amount of organic solvent added impacted the integrity of the plasma samples over time. The ion suppression observed in plasma was not due to the addition of the CAD (naltrexone) but due to the amount of organic.

521 Dried Blood Spots (DBS) On-Card Derivatization: An Easy & Alternative Form for Sample Handling to Overcome the Biological Matrix Instability of Thiophinan.


Purpose: Use of in-house pre-treatment of DBS cards to overcome stability challenges of thiophinan in toxicokinetic and pharmacokinetic studies.

Methods: Thiophinan was spiked in whole blood (range of 5-600 ng/mL) and 40 ul of whole blood were applied on untreated or in-house pretreated cards with 2-bromo-3'-methoxycatecholone FTA DMPK-A, B and C cards. Disks (6 mm) were punched from the blood spots, transferred into tubes and 200 ul of internal standard (thiorphan-D5) in methanol was added. Samples were vortexed and left on bench for one hour. 100 ul of the supernatant was transferred into 96 well plate already containing 100 ul of water prior to injection on a Zorbax Bonus RP column eluted using a gradient. Analysis was performed on an ABSciex API3000 in electrospray (+).

Results: The instability of thiophinan in biological matrix was demonstrated and minimized by derivatization to thiophinan-MP with 2-bromo-3'-methoxycatecholone (BMP). In order to simplify sample handling process, on-card derivatization using DBS was investigated. The instability of thiophinan on-card, without derivatization, was demonstrated on all DBS cards used. Therefore, on-card derivatization technique was investigated. BMP could be added directly to the card box at least 2 days prior to blood spotting without compromising thiophinan derivatization. Although the on-card derivatization of thiophinan was successful on the A and C cards, the derivatization reaction seemed to be inhibited on the B card. Linearity of a calibration curve for a range of 5-600 ng/mL was demonstrated using A and C cards pretreated with 0.5M BMP. Precision was 2.8% and the accuracy was 97.104%. Results showed on-card stability of thiophinan-MP for at least 2 days.

Conclusion: It was clearly demonstrated that DBS on-card derivatization offers a practical alternative for the bioanalysis of thiophinan in TK & PK studies since it requires less manipulation than the regular sample handling process.

522 Superparamagnetic Iron Oxide Loaded Cross-Linked Nanosamplons Improve Tumor Accumulation and Magnetic Resonance Imaging In Vivo.

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Purpose: To improve accumulation and contrast-enhanced magnetic resonance imaging of glioma using cross-linked nanosamplons loaded with superparamagnetic iron oxide nanoparticles (CNA-IONPs) for potential glioma therapeutics.

Methods: CNA-IONPs were synthesized and characterized. Gd-DTPA was used to monitor glioma tumor growth as a T1 contrast agent. CNA-IONPs were injected via the tail vein of a glioma xenograft rat model. T2 weighted MR images were taken via the tail vein of a glioma xenograft rat model. T2* was used to predict CNA-IONPs concentration increases. Glioma tumor, contralateral brain tissue, blood, and peripheral organs were collected at 2 or 6 h and iron concentrations analyzed inductively coupled plasma mass spectrometry (ICP-MS).

Results: CNA-IONPs had a neutral zeta potential, hydrodynamic size = 30 nm, and were highly stable in medium containing serum at 37 °C over 30 h. CNA-IONPs showed significant T2 and T2* contrast enhancement 2 h after injection. Gd-DTPA (0.3 mmol/kg) enhanced the tumor on T1 images acquired immediately after injection. T1 images acquired 2 h after injection showed Gd-DTPA was almost entirely cleared. In certain regions of the tumor, R2* (1/T2*) was proportional to iron concentration, increased 48% from 10 min to 2 h after CNA-IONP injection. ICP-MS iron concentration in glioma tissue was 3 to 9 times higher than in contralateral brain tissue at 2 h. The iron concentrations in blood and peripheral organs did not significantly change from 2 to 6 h.

Conclusions: CNA-IONPs had desirable physicochemical properties for in vivo application. They preferentially accumulated in glioma and the concentration increased with time consistent with their long blood circulation time. CNA-IONPs have the potential to improve MRI diagnosis and provide a platform for incorporating glioma therapeutics.

523 Age-Dependent Capability of Drug Metabolism in Commercially Available Human Hepatocytes.

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Primary human hepatocytes have been widely used as a good tool to evaluate the safety of chemicals containing environmental pollutants, pesticides and drugs.

On the other hand, it is known that the drug-metabolizing enzyme activities of hepatocytes are changing with aging. There, however, are large interindividual variations in the levels of CYP enzyme activities and the response to inducers among hepatocyte donors. Therefore, in this study we analyze the relation between the enzyme activities and the induction levels of the enzymes.

We analyzed the enzyme activities of CYP1A1, CYP2B6, CYP3A4, CYP2C8, CYP2C9, CYP2C19, CYP2D6 and UGT and their responses to CYP inducers by comparing the data of cryopreserved human hepatic cell suspension and plateable hepatocytes regarding the enzyme activities and the levels of mRNAs encoding the enzymes presented from multiple suppliers. Total samples used in this study were obtained from more than 100 donors (5 months - 82 years old).

CYP1A2 activities in hepatocytes from young donors (< 4 years old) tended to be reduced in cell suspension and were largely induced both in the activity and the mRNA levels in plateable hepatocytes by beta-naphthoflavone. The enzyme activities and the mRNA levels of CYP3A4 and CYP2B6 in the young donors were positively induced by rifampicin. There was large variability in CYP activities and the abilities of the inducers. The utility of these cryopreserved human hepatocytes as an evaluation system for the safety of chemicals will be discussed.

524 AhR-Mediated Epigenetic Modulation of CYP19 and BCRP in Human Breast Adipose Fibroblasts and MCF-7 Cells.

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The epithelial-stromal microenvironment in breast cancer largely determines chances of therapeutic success, due to modulation of tumor cell gene expression and drug resistance. In breast adipose fibroblasts (BAFs) surrounding a tumor, expression of aromatase (CYP19), the enzyme responsible for local estrogen production, is elevated. In breast cancer cells, over-expression of breast cancer resistance protein (BCRP) is one of the major causes of multi-drug resistance and chemotherapeutic failure. Both BCRP and CYP19 over-expression have been related to hypomethylation of the promoter regions by DNA methyltransferases (DNMTs).

Earlier studies have suggested that the aryl hydrocarbon receptor (AhR) can function as an epigenetic regulator. Here, the effect of the non-toxic AhR agonist tramilast on CYP19 and BCRP expression was investigated in BAFs and MCF-7 cells. In BAFs, tramilast caused a significant 70% reduction of CYP19 expression, which concurred with a 1.3/pII to 1.4-promoter switch. In MCF-7 cells, tramilast induced expression of the AhR-responsive gene CYP1A1 (5-fold) and breast cancer resistance protein (BCRP, 2-fold), but had no effect on the expression of interindividual variability. CYP1A2 activities in hepatocytes from young donors (< 4 years old) tended to be reduced in cell suspension and were largely induced both in the activity and the mRNA levels in plateable hepatocytes by beta-naphthoflavone. The enzyme activities and the mRNA levels of CYP3A4 and CYP2B6 in the young donors were positively induced by rifampicin. There was large variability in CYP activities and the abilities of the inducers. The utility of these cryopreserved human hepatocytes as an evaluation system for the safety of chemicals will be discussed.

525 The Ah Receptor Recruits Protein Kinases to Phosphorylate Ser10 in Histone H3 of the CYP1A1 Promoter Chromatin.

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Halogenated aromatic hydrocarbons such as 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) cause many types of toxicity via the aryl hydrocarbon receptor (AhR). One of the best known proteins induced by TCDD is the cytochrome P450 1A1
methylation patterns are similar between the age groups, but there are dis-
tinct methylation patterns between male and female rats. The results indicate that
there are slightly more methylated sites at 2wk of age compared to later ages. A
higher number of methylated sites were found on the X chromosome in females
than males, suggesting involvement in repression of X-inactivated genes. These
results provide a comprehensive global view of the DNA methylation status in
the kidney over the entire rat life cycle. These age- and sex-related differences in DNA
methylation may provide insights in susceptibility to kidney disease and its progres-

528 Phylogenetic Identification of Variably Methylated Transposons As Biomarkers for Early Environmental Exposures.
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A small number of retrotransposons in the long terminal repeat (LTR) class are
known to exhibit environmental sensitivity in DNA methylation status, and are
termed “metastable epialleles”. Two of the sequenced examples, the agouti viable
yellow (AY) and the CDK5 activator-binding protein (CbpAP) result from a re-
cently inserted LTR, an intracisternal A particle (IAP) element. These insertions are
from the same family, and show high sequence identity (98.5%). Additionally, both
act as biomarkers of early exposure. Until now, neither the full extent of variability
at these metastable epialleles, nor the phylogenetic relationship underlying variable
elements was well understood. Using a computational approach, we identified
10,802 IAP LTRs in mice, and filtered by subtype to yield 1,388 IAP LTRs in the
family that includes AY and CbpAP. Phylogenetic analysis revealed duplication
and divergence events subdivide this family into three clades. We characterized
variability across clades, DNA from isogenic mice was subjected to combined bisulfit
and restriction analysis (CoBRA) at 21 LTR transposons (7 per clade). To validate
our candidate IAP LTRs for interindividual variation, we assayed 17 isogenic mice
for shifts in liver DNA methylation patterns. Methylation levels at individual LTRs
varied widely with mean methylation ranging from a low of 59% to a high of 89%.
Among mice, average methylation across all LTRs was not significantly different
(71%-74%, p > 0.99). Finally we determined that the clade with the most con-
served elements had significantly higher average methylation across LTRs than ei-
ther diverged clade. Thus, we increase the number of known epigenetically modifi-
able loci and provide evidence that sequence identity is predictive of methylation
level. Since repetitive elements comprise nearly half of mammalian genomes, they
are likely targets for toxicological disruption, especially during early development.
The characterization of murine metastable epialleles is crucial for the development
of treatments for environmentally-induced disease.

529 Chromatin Context Modules Genomewide p53 Sequence-Specific Occupancy and Exposure-Induced Gene Expression.
DNA damaging agents activate p53 to bind to DNA response elements leading to
transactivation of p53 pathway genes and directing cells toward arrest or apoptosis.
To understand the determinants of treatment-specific and tissue-specific responses to
DNA damage, we have carried out treatments using ENCODE cells in an expos-
ure model and generated genomewide data (p53 ChIPseq, H3K4me3 ChIPseq,
sequence specificity, evolutionary conservation and gene expression). Effects of
doxorubicin and nutlin-3 treatments in lymphoid cells have been compared with
other cell types and analyzed in relation to chromatin context based on the EN-
CODE chromatin hidden Markov model (ChromHMM). We identified novel
chromatin interactions that modulate the transcriptional response to p53 activation
following DNA damage. Highly induced p53 genes typically display both low
H3K4me3 marks and low baseline expression. Among the 2568 high confidence
p53-occupied genes detected by p53 ChIP-seq, 45% have active promoter marks
(H3K4me2, H3K4me3, H3K27ac, H3K9ac) prior to treatment. However, surpris-
ly, based on ChromHMM categorization, ~22% of p53 genes displayed repressive
chromatin states; including many highly-induced p53 response genes. Many p53
peaks were located in genomic regions with preexisting states classified as promot-
ers (16%) or enhancers. However, notably a large percentage (31%) of all p53 bind-
ing peaks were found in repressed or heterochromatin regions far distant from a
transcription start site. Thus p53 binding occurs in regions of both accessible and
inaccessible chromatin although induced expression was most commonly associ-
ated with inaccessible chromatin. We identified p53 DNA binding motifs in 95% of
p53 ChIPseq peaks and used a position weight matrix model (PWM) to score
similarity to the consensus p53 motif. PWM strength and spacer length have dis-
tinctive distribution patterns between different chromatin states. Thus both epi-
genetic and sequence-based factors modulate the DNA damage response regulated by
p53.
J. Goodrich, N. Bau, A. Framblau and D. Dolinoy, Environmental Health Sciences, University of Michigan, Ann Arbor, MI.

Epigenetic alteration may be a key mechanism linking chemical exposures to toxicity and disease. Recent, yet limited, animal and human data suggest that mercury (Hg) modifies the genome, specifically DNA methylation. This study hypothesizes that methylmercury and inorganic Hg exposures from fish consumption and dental amalgams, respectively, alter DNA methylation patterns at multi-copy repeats and candidate genes. Dental professionals were recruited at meetings of both the Michigan and American Dental Associations (MDA, ADA). Subjects provided survey data (e.g. exposure sources, demographics) and samples for Hg measurement and epigenetic analysis. Total Hg was quantified via atomic absorption spectrophotometry in hair and urine, indicative of methylmercury and inorganic Hg exposures, respectively. Methylation was assayed globally by pyrosequencing of long interspersed elements (LINE-1) and site-specifically at Dnmt1, Seppl1, and Seppl2 in bisulfite converted DNA (isolated from buccal mucosa or saliva in the ADA). In the MDA cohort (n=158), hair Hg (mean ±SD: 0.55 ±0.54 µg/g) and urine Hg (1.05 ± 0.81 µg/L) overlapped with levels of the general US population. Multivariate linear regression displayed a trend of hypomethylation at LINE-1 and three candidate genes with increasing Hg levels. Only in males, this trend was significant (p<0.05) in models of mean blood leukocytes and saliva in the ADA). In the MDA cohort (n=158), hair Hg and Dnmt1 methylation were not significant. Associations between Hg biomarker levels and DNA methylation will be further explored using samples from the ADA cohort (n=500). This work suggests that methylmercury modifies DNA methylation at labile regions in males. Epigenetic modification by Hg should be further investigated to elucidate this pathway as a mechanism of toxicity.

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Blueberry (Vaccinium spp.) has shown a broad spectrum of biomedical functions including anti-oxidative and anti-inflammatory effects through in vitro or animal studies. However, its human studies are relatively few. In addition, food-born epigenetic modulators, e.g. folate, genistein, etc., can recover environmental exposure -related epigenetic alterations. Therefore, epigenetics can be a useful screening method to find a new functional food. We performed a two weeks intervention study to evaluate anti-oxidative and epigenetic effects of blueberry in young women (N=8, age 22-60±0.7 yrs, BMI=20.2±1.97 kg/m²). Among the subjects, two persons took vt C (1 g/day) as positive controls and the others took blueberry juice (240mI total polyphenol 300mg, anthocyanin 76mg/day). From genomic DNA of their peripheral blood, we analyzed global methylation-status with methylFlashTM methylated DNA quantitation kit (Epigenetix). We also analyzed urinary malondialdehyde (MDA), a biomarker of oxidative stress with HPLC/UVD. As a result, we found that blueberry did not reduce urinary MDA levels as much as vt C. Additional work is needed to find out what increased global DNA methylation. There were no significant differences in the global DNA methylation between the two treatments (p=0.89). Therefore, anti-oxidative activity of blueberry was not found in this trial, compared to vt C. However, its epigenetic effects can be similar to vt C’s. In conclusion, the present study provides that a medicinal function of blueberry can be induced by global hypermethylation rather than anti-oxidative effects.

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This study sought to investigate the role of microRNAs (miRNAs) in the pathogenesis of MMPI-induced skin fibro dysplasia. MicroRNAs are short non-protein-coding RNAs that modulate protein translation from specific mRNAs. Certain miRNAs exhibit tissue specificity, and are dysregulated in response to specific pathologies. MicroRNAs are detectable in biological fluids, paving the way for their use as biomarkers. Matrix metalloproteinases (MMPs) are a family of proteolytic enzymes and have been an attractive pharmacological target for a number of indications. However, development has been hampered by the propensity of these compounds to cause connective tissue pathologies. The broad-spectrum MMP inhibitor AZM551248 has been shown to induce such effects in the dog, characterised by fibroblast proliferation and collagen deposition in subcutaneous tissues. Thirty 12-month old female beagle dogs were assigned to six groups. Animals were dosed orally once daily with vehicle, or with vehicle plus 20mg/kg/day AZM551248 for between 4 and 17 days. miRNA expression profiles in subcutaneous fluids, the primary target of MMPI skin, were determined by microarray analysis. miRNA expression profiles in subcutaneous fluids of MMPI-treated dogs were compared to healthy controls. Following 4 days MMPI administration, 13 miRNAs were differentially expressed in the skin compared with vehicle treated animals. Several of these members of the miR-200 family were attenuated in response to MMPI. As the severity of FD increased at the later time-points, miRNAs associated with TGFβ synthesis were upregulated. Evidence of epithelial to mesenchymal transition was present at all study time points. Receiver operator curve analysis revealed that miR-21 expression in the cervical subcutaneous tissue was a sensitive and specific biomarker of FD in cidence.
We have described a role for miRNAs in processes relevant to the key histopathological events associated with MMPI-induced FD. Furthermore, we have identified key miRNAs with the potential to be used as informative biomarkers of FD.

535 Exposure of Pregnant Mice to Chlorpyrifos-Methyl Alters Embryonic H19 Gene Methylation Patterns.

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The aim of this study was to identify whether chlorpyrifos methyl (CPM) exposure during pregnancy leads to changes in the methylation patterns of H19 gene. CPM 4, 20, 100mg/kg bw/day was administered to 4 pregnant mice per group between 7 and 12 days post coitum (d.p.c.). Pregnant mice were killed at 13 d.p.c. and genomic methylation in primordial germ cells (PGCs) and fetal organs (the liver, intestine, and placenta) was examined. Four polymorphism sites in the H19 alleles of maternal (C57BL/6J) and paternal (CAST/Ei) alleles were identified at nucleotide position 1407, 1485, 1566 and 1654 and the methylation patterns of 17 CpG sites were analyzed. The methylation level in male and female PGCs was not altered by CPM treatment in the maternal allele H19. The methylation level of the paternal H19 allele was altered in only male PGCs in response to the CPM treatment. The methylation level at a binding site for the transcriptional regulator CTCF2 was higher than that at the CTCF1 binding site in all CPM-treated groups. In the placenta, the aggregate methylation level of H19 was 56.89% (control) and ranged from 47.7% to 49.89% after treatment with increasing doses of CPM, H19 gene from thelifer and intestine of 13 d.p.c. fetuses treated with CPM was hypomethylated compared with controls, although H19 mRNA expression was unaltered. In the placenta, H19 expression was slightly increased in the CPM-treated group, although not significantly. IGF2 expression levels were not significantly changed in the placenta.

In conclusion, CPM exposure during pregnancy alters the methylation status of the H19 gene in PGCs and embryonic tissues. We infer that these alterations are likely related to changes in DNA demethylase activity.

536 Autoimmune Disease Triggered by Trichloroethylene Is Associated with Epigenetic Alterations inCd4+ T Cells.

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Previous studies have shown that chronic (32-week) exposure to occupationally relevant concentrations of trichloroethylene (TCE) in the drinking water of female MRL/lpr mice promoted autoimmune hepatitis. This was accompanied by the expansion of CD4+ T cells that secreted increased levels of IFN-γ and expressed an activated (CD44+CD26+) phenotype. The current study was initiated to determine the mechanism by which TCE altered CD4+ T cell function. The study conducted a longitudinal evaluation of mouse TCE exposure over 40 weeks. Alterations in IFN-γ production corresponded to changes in the expression of markers used to assess DNA methylation, namely retrotransposons Iap (Intracisternal A Particle) and MuERV (murine endogenous retrovirus). In addition, global DNA methylation was significantly altered in CD4+ T cells from TCE-treated mice. Most recently, bisulfitie sequencing revealed that DNA methylation of Cpg sites associated with the Ifng promoter was significantly, and time-dependently altered by TCE exposure.

Placentation of epigenetic effects appropriately into the product safety paradigm is key miRNAs with the potential to be used as informative biomarkers of FD. In conclusion, CPM exposure during pregnancy alters the methylation status of the H19 gene in PGCs and embryonic tissues. We infer that these alterations are likely related to changes in DNA demethylase activity.

537 Is the Current Product Safety Assessment Paradigm Protective for Transgenerational Epigenetic Effects?

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Placement of epigenetic effects appropriately into the product safety paradigm is challenging due to an incomplete understanding of causal associations between epigenetics and apical adverse effects. To begin to understand this relationship, groups of 25 pregnant F0 CD-1 mice were administered -10 μg/kg/day diethylstilbestrol (DES) or -30 μg/kg/day 17β-estradiol (E2) via the diet or by subcutaneous injection (SC) from gestation day 9 – lactation day 20. F1 offspring were mated for two additional generations, with F1-F3 offspring evaluated for uterotropic effects on postnatal day (PND) 21 and 18 months. F1 generation females in the DES and E2 diet groups had a 3-4-fold increase in relative uterine weights, which corresponded to plasmacytosis and hypertrophy of the endometrium and squamous cell carcinoma in the DES diet group, but no histopathological effects in either E2 group at 18 months. When comparing gene lists in the PND 21 E2 and DES diet groups, ~1500 known genes were unique for DES, of which 24 remained significantly altered up to 18 months. These genes sets provided candidate markers for future epigenetic analyses. F2 generation females in the DES diet group had a marginal increase in mean PND 21 uterine weights and a treatment-related increase in leiomyoma/leiomyosarcoma at 18 months. In the F3 generation, there was no effect on uterine weight or histopathology at PND 21 or 18 months. Under the conditions of this study, there were no transgenerational effects associated with DES or E2, which would suggest that a typical margin of exposure risk assessment would be protective.

538 Genome-Wide Methylation Profiling in Rat Tissues for Nongenotoxic Carcinogens Using Medip-Chip.

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DNA methylation is an epigenetic mark that has a major role in carcinogenesis. The aim of the study was to perform a genome-wide analysis of DNA methylation profile of non-genotoxic hepatocarcinogens including tetrachlorodibenzo-p-dioxin (TCDD), hexachlorobenzene (HCB), methyrapline (MYP) and the male rat kidney carcinogens, d-limonene, p-dichlorobenzene (DCB), chloroform and ochratoxin A (OTA) in their respective target tissues in rats dosed under bioassay conditions. Methylated DNA immunoprecipitation (MeDIP) was used in conjunction with a NimbleGen promoter plus CpG island array to identify differentially methylated DNA regions (DMRs) on a genome-wide basis. We first identified a total of 76 candidate DMRs for OTA, 1 DMR for MYP and 4 DMRs for chloroform by the MeDIP-Chip method. No change was observed in rats treated by TCDD, HCB, DCB and d-limonene. It is expected that results from this study will contribute to a better understanding of the key molecular events which occur during early stages of chemical carcinogenesis and aid in identification of mechanism-based biomarkers for early detection of non-genotoxic carcinogens.

539 Modulation of DNA (Cytosine-5)- Methylation in the Mouse Embryonic Stem Es-D3 Cell- and in the Human Intestinal CaCo2-Cells by Okadic Acid.

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Okadic acid (OA) a polyether fatty acid produced by marine plankton (dinoflagellates) accumulates in the black sponge and in mussels and oysters and causes shellfish diarrheic syndrome in consumers. It is also suspected of causing gut tumours in humans. It is a tumour promoter in mouse skin, inhibitor of protein phosphatase 1 (PP1) activity, it should be taken into account when assessing the risk for human.

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Previous studies from our laboratory demonstrated that expression of the human epoxide hydrase gene (EPHX1) is driven by a far upstream E1-b promoter in most human tissues, whereas the proximal E1 promoter selectively drives hepatic EPHX1 expression. The molecular mechanisms underlying the differential promoter usage are not well elucidated. CpG islands are short stretches of DNA harboring relatively higher frequencies of the cytosine/guanine repeat sequence, potentially reflective of higher gene methylation modifications. In-silico analysis of the human EPHX1 gene indicated that the upstream regions of E1 and E1b promoters of human EPHX1 gene are both rich in CpG islands. As methylation status of CpG islands around gene promoter regions is often associated with transcriptional repression of genes, we investigated the methylation status of CpG islands proximal to the upstream regions of the E1b and E1 promoters in several human cell lines and in primary human hepatocytes. Our data demonstrate that the methylation status of all the CpG islands in upstream regions of the E1b promoter from different cell lines is essentially the same and only sparsely methylated (<10%). In contrast, the CpG islands located at regions close to the E1 promoter are highly methylated (>60%) in cell lines originating from lung and kidney. In contrast, in hepatocellular carcinoma HepG2/C3A cells, the CpG islands localized at -128bp, -89bp, -488bp and +8bp of the E1 promoter are 0%, 0%, 0%, 0% and 25% methylated, respectively. Similarly, the CpG islands located at -724bp, -602bp, -128bp, -89bp, -488bp and +8bp of the E1 promoter in human primary hepatocytes are all unmethylated. The methylation status of these promoters from human hepatic and non-hepatic tissues are also consistent with our results from different cell lines and primary hepatocytes. These findings suggest that differential methylation status determines tissue selective promoter usage for EPHX1 expression in humans.

541 Identification of Arsenic-Responsive microRNAs in Rats by Genome-Wide High-Throughput Sequencing.

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Consumption of drinking water contaminated with arsenic, a naturally occurring carcinogenic metalloid, constitutes a major public health problem. Although the relationship between exposure and carcinogenesis is well documented, the mechanisms by which arsenic participates in tumorigenesis are not fully elucidated. Epigenetic modifications, other epigenetic mechanisms, in particular miRNA expression, that play a critical role in regulation of gene expression, have yet to be adequately investigated for arsenic. In the current study, all rats were exposed to sodium arsenite drinking water for 60 days. miRNAs was extracted from rat liver, the primary tissue for arsenic metabolism and potential target of arsenic carcinogenesis, following As(3+) treatment. Genome-wide profiling of miRNA in rat livers was generated to identify As(3+)-responsive miRNAs by next generation high-throughput sequencing technology. The preliminary results indicate that exposure to environmentally-relevant levels of As(3+) lead to aberrant expression of multiple miRNAs in rat liver. Additional analysis will identify miRNAs whose levels are most differentially expressed in rats due to arsenic treatment, and potentially identify early pre-disease epigenetic biomarkers of arsenic exposure. The results may also provide insights into the role of miRNAs in arsenic carcinogenesis (This work was supported by Zhejiang Natural Science Foundation, China (H.W.); and Startup fund (to X.R.) provide by SUNY at Buffalo).

542 Benzo(a)pyrene Enhances Line-1 Reactivation in the Absence of Retinoblastoma Proteins: Implications for Global Epigenetic Silencing and Heterochromatin Regulation.

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Benzo(a)pyrene (BaP) is a ubiquitous environmental pollutant that induces DNA mutations and disease. At the molecular level, BaP induces the reactivation of LINE-1 (or L1), a primate-specific transposable element that normally acts as a silenced, differentiated cell. L1 epigenetic silencing involves promoter DNA hypermethylation, re-activation of transcriptional repressors such as retinoblastoma (RB), and small RNAs. Since RB proteins control the cell cycle and help recruit transcriptional co-repressors to chromatin, we hypothesized that i) L1 expression may follow cell cycle progression, and ii) absence of RB is associated with derepression of L1. In the present studies, ES-D3 mouse embryonic stem cells were stained live and sorted for different phases of the cell cycle, followed by measurements of L1 message by quantitative PCR. L1 message was most abundant in the G1 phase of the cell cycle, coinciding with the stage where L1 became activated by hypomethylation. In separate experiments, isolates of mouse embryo fibroblast (MEFs) deficient for the RB family of proteins (TKO MEFs) were exposed to BaP for 12 hours and real time PCR used to quantify changes in the abundance of L1 transcripts. Compared to wild type counterparts, RB null cells dramatically overexpressed L1 in the presence of BaP. Cytochrome P450 genes involved in BaP metabolism also showed an enhanced inducibility in mutant cells, suggesting increased chromatin accessibility to transcription factors. Collectively, these data indicate that L1 is regulated in a cell-cycle dependent manner and that mutation or loss of RB proteins in humans in the face of environmental stress may contribute to the deregulation of L1 in somatic cells. Such deregulation may compromise differentiation programs and induce acquisition of neoplastic phenotypes.

543 DNA Methylation Patterns Represent “Environmental Footprints” of Transcription Factor Occupancy.

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Previous research has demonstrated that binding of transcription factors (TFs) to promoters can alter the DNA methylation patterns within these regions. While previous research was conducted in the context of stem cell differentiation, the impact of environmental toxicants on TF binding and subsequent effects on DNA methylation has not been examined. Our group recently identified genes in human leukocyte DNA and mouse livers that exhibit differential CpG island methylation following exposure to metals/metalloids including cadmium and arsenic. Among these genes, we have identified enrichment for transcription factor binding sites that we postulate represent “environmental footprints” of transcription factor occupancy. Based on these results, we hypothesize that in response to environmental stimuli, selective occupancy of DNA regions by transcription factors can alter the access of epigenetic modifiers to DNA, resulting in gene-specific DNA methylation patterns. To support our hypothesis, we utilized in vitro culture of human cells to study changes in genome-wide promoter DNA methylation patterns associated with specific changes in transcription factor binding induced by cadmium and arsenic exposure. Specific genes were identified among these that showed up-regulation in response to each metal by which specific genes display toxicant induced patterns of DNA methylation.

544 Increased Growth Inhibition of Human Breast Cancer Cells by Co-Treatment with EGCG and 5-aza-2-deoxycytidine Is Modulated through Epigenetic Mechanisms.

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Breast cancer is the leading cause of cancer deaths in women. Both genetic and epigenetic processes are implicated in breast carcinogenesis. Alterations in DNA methylation pattern contribute to cancer development. Studies show that half of the tumor suppressor genes are inactivated more often by epigenetic, rather than genetic mechanisms in all types of cancer. As epigenetic changes like DNA methylation are reversible, demethylating drugs, such as, 5-aza -2’ deoxycytidine (5-aza) are being clinically used as cancer chemotherapeutics. However, there are cytotoxic effects associated with high doses of these drugs. Alternative dietary compounds like EGCG, a green tea polyphenol known to possess anticancer properties with minimal or no toxic effects on normal cells. EGCG is also reported to have DNA demethylation and histone modification activity. Therefore, the purpose of this study was to investigate the efficacy of co-treatment of EGCG and 5-aza, at low doses in inhibiting growth of MCF-7 breast cancer cells. MCF-7 cells were treated with either 5-aza, (5μM), EGCG (50μM, 100μM) or their combination. MTT assay showed that co-treatment significantly decreased cell viability as compared to
individual treatment. Both compounds arrest separate phases of cell cycle, but co-
treatment individually increased the percentage of growth arrested and apoptosis in cells. Increased hypomethylation due to co-treatment was also observed in global genomic methylation analysis. Downregulation of DNMT1 expression was significantly greater in co-treatment than individually treated cells. These data suggest that co-treatment with EGC2 and 5-aza, at low doses can synergistically reverse hypermethylation induced gene silencing and hence can be used for inhibition of breast cancer growth while minimizing cytotoxicity on normal cells.

545 Differential Expression of Long Intervening Noncoding RNAs in the Livers of Female B6C3F1 Mice Exposed to the Carcinogen Furan.

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The mammalian genome is transcribed into mRNAs that code for protein as well as a broad spectrum of other noncoding (nc) RNA products. Long ncRNAs (lncRNA), defined as noncoding RNA species > 200 nucleotides long, are emerging as important epigenetic regulators of gene expression and are involved in a spectrum of biological processes of relevance to toxicology. We have conducted a gene expression profiling study in liver using female B6C3F1 mice exposed to the carcinogen furan at 0.0, 1.0, 2.0, 4.0, and 8.0 mg/kg for 3 weeks. lncRNAs showed a non-linear dose response with no lncRNAs differentially expressed at 1.0 or 2.0 mg/kg and two lncRNAs differentially expressed at 4.0 mg/kg furan. At 8.0 mg/kg furan, 13.3% (83/632) of the differentially expressed transcriptome was comprised of lncRNAs. Among the lncRNAs observed, a number showed transcriptional clustering with nearby protein coding genes. For example, in furan-exposed mouse liver there was increased expression of lncRNA-p21. LncRNA-p21 is an anti-sense transcript that is 15 kb downstream from Cdkn1a locus and is known to be induced by p53 in human cells. LncRNA-p21 appeared to be co-transcribed with the protein coding gene Cdkn1a in response to 8.0 mg/kg furan. These data suggest that lncRNAs are transcriptional targets of furan-induced cytotoxicity and cell proliferation. We hypothesize that lncRNAs have potential as epigenomic biomarkers of carcinogenic exposures and we are currently examining lncRNA expression in the liver of mice exposed to a number of other carcinogens.

546 ROS-Generation Leads to 5-Hydroxymethylcytosine Formation and DNA Methylation.

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DNA methylation is a chemical modification at the 5' position on cytosine residues involved in gene repression. Notably, DNA methylation plays a vital role in stem cell differentiation. A better understanding of the dynamic processes controlling methylation and demethylation is critical to identifying chemicals with potential to disrupt programming of the methylome and may uncover targets in diseases such as neurodegenerative disease and cancer. While enzymes which methylate cytosines neighboring guanines (CpGs) have been identified, no demethylase has been discovered in mammalian cells. Inhibition of DNA methyltransferases leads to demethylation, though these mechanisms depend on cell division and may not explain active demethylation in post-mitotic cells such as neurons. Recent work has uncovered the role of Tet proteins in converting 5-methylcytosine to 5-hydroxymethylcytosine (5-hmC). Subsequent deamination by cytidine deaminases followed by base excision repair results in unmethylated cytosine. Here, we show that sub-lethal exposures of the benzene metabolite hydroquinone (HQ) led to formation of 5-hmC in HEK293 cells and a human neural progenitor cell line. In HEK293 cells, HQ exposure led to increases in reactive oxygen species (ROS), re-activation of a methylation-silenced reporter plasmid, and demethylation of CpGs within its promoter. Both 5-hmC formation and plasmid reactivation were inhibited by the antioxidants α-tocopherol and N-acetyl-L-cysteine, indicating ROS formation was involved. Moreover, reactivation of the silenced plasmid by HQ was enhanced in cells overexpressing cytidine deaminase APOBEC2. These effects were observed with no detectable changes in Tet gene expression, though activity may have changed. Our data indicate toxicant-induced ROS leads to CpG demethylation in a manner consistent with the demethylation-deamination mechanisms catalyzed by Tet proteins and suggest a link between cellular redox status and maintenance of the epigenome.

547 A Critical Review of Epigenetic Transgenerational Inheritance following Environmental Exposures.


Epigenetic transgenerational effects are phenotypic traits passed down from generation to generation via heritable modifications in the epigenetic landscape. There are several accounts in the literature investigating the effects of environmental factors on the transgenerational inheritance of purportedly epigenetic traits. Epigenetic transgenerational effects have been suggested through animal models to lead to such conditions as polycystic ovarian disease, prostate disease phenotype, abnormal sperm parameters, altered discrete brain nuclei, fetal abnormalities, increased tumor frequency, altered immune function, and indications of kidney disease. Certain compounds have received particular attention with respect to transgenerational epigenetic changes, with some publications claiming to demonstrate transgenerational epigenetic effects and others refuting the existence of these effects. Recently, increased interest in these diverse compounds has been tested for the ability to induce transgenerational epigenetic changes either alone or in mixtures, including pesticides, plastics components, and hydrocarbons. Although many publications claim to demonstrate clear evidence of transgenerational epigenetic effects, an equally clear mechanism explaining how the effect is conveyed is often lacking. Additionally, it remains to be explained how any particular chemical, that is not, for example, a methyl donor, could impact such a diverse collection of phenotypic outcomes. While the possibility of epigenetic transgenerational inheritance is not questioned, it is suggested that a clear or strong mechanistic link has not been established between many of the compounds purported to effect epigenetic transgenerational inheritance, and the subsequently observed phenotypes. A critical analysis of a selection of the evidence assessing epigenetic transgenerational inheritance following exposure to environmental agents and chemicals will be presented with particular focus on mechanisms of epigenetic inheritance, discrepancies in the literature, study quality, reproducibility and transparency.

548 Effect of Increased Reactive Oxygen Species on microRNA Expression in First Trimester Placental Trophoblasts.

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In early pregnancy, the conceptus is highly sensitive to reactive oxygen species (ROS). Expression of superoxide dismutase, catalase, glutathione peroxidase and high levels of glutathione, taurine, and vitamins A and D protect the conceptus by 5 weeks. During implantation, maternal spiral artery remodeling and trophoblast invasion serves as the beginning of placental development and would result in increased protection of the fetus to normal ROS signaling. However, improper invasion and remodeling can result in disease development. Although debated, the current proposed model for preeclampsia, which manifest as abnormally high blood pressure (>140/80) after 20 weeks gestation in a previously normotensive woman, is aberrant maternal spiral artery remodeling, subsequent altered angiogenesis and localized ischemia/reperfusion due to continued muscularization of the spiral arteries. Preeclampsia is associated with low birth weight, preterm labor, small gestational age, and intraterrine growth restriction as well as altered gene expression in the placenta as a result of increased ROS production. Epidemiological screening studies for preeclampsia indicate alterations in placental microRNA (miRNA) expression as a result of increased trophoblast apoptosis. The goal of this project is to determine the effect of increased ROS during early placental development on miRNA expression to better understand the time frame of gene expression alteration. The villous 3A first trimester cell line was exposed to varying concentrations of H2O2 over 24 hours to determine the amount of intracellular ROS generated from this model. Using an optimal exposure of 50 μM, we then determined the effect of increased ROS exposure on the expression of 1000 miRNAs using the human microRNA PCR array (Qiagen). The miRNAs shown to be altered from increased ROS production are being selected for pathway analysis and further study using miRNA mimics and inhibitors (Supported by T32-07454; P30-006676).

549 Identification of Mode-of-Action Specific Toxicity Transcript Profiles In Vitro Using a Connectivity Mapping Approach.

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We hypothesize that it is possible to predict toxicity by assessing changes in gene expression in a small number of cell types enriched in toxicologically relevant pathway and determined the connectivity profile of 34 chemicals in HepG2, Ishikawa and MCF7 cells, using microarrays (U219, Affymetrix). These chemicals act via a number of modes of toxicity and do not involve direct chemical
reactivity, as this can be adequately detected by a number of simple in vitro methods. Each of the chemicals was tested at 3 doses and at 3 different time points; however, data presented here are for only one dose and one time point (6h). The relatedness of each transcript profile to each other was analyzed using the cMAP platform (Broad Institute). The results indicate that not all cell lines responded to all chemicals. For example the response of the Ishikawa cells to methotrexate is minimal (16 genes significantly affected), while the MCF7 and HepG2 cells have a robust response to this chemical (>3000 genes significantly affected). A comparison of responses to chemicals that share similar modes of action (MOA) exhibited very high correlation, even among different cell lines. These correlations held true for both agonists (cMAP score ≥ 1) and antagonists (score ≤ -1) and extended to MOAs mediated through receptors (AR, ER, FX, PR, etc.) and enzyme inhibition (histone deacetylase, DHFR, etc.). For example, using the signature from valproic acid (VA) treated cells to query the cMAP database yield high positive connectivity scores for VA in all three cell lines tested, as well as with other HDAC inhibitors (vorinostat, trichostatin, butyoxac, depudac and HC toxin), regardless of cell lines or concentrations. The transcript profile elicited by chemicals with various MOAs in selected cell types, coupled with a cMAP approach for the analysis of the response, offers relevant biological data to predict biological activity and MOA, and thus toxicity, in a defined in vitro system.

550 Analysis of Longitudinal Metabolomic Data from Endocrine Disruption Studies: The A-SCA Method.

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Metabolomics are increasingly being used in the field of toxicology: Experimental designs involving the study of dynamic changes in the metabolome raise new methodological challenges in the field of data analysis, regarding long term and perinatal in vivo studies. Multivariate analysis of variance (MANOVA), often used to analyze experimental data, is not always appropriate for metabolomics, especially when sample size is much smaller than the total number of variables, which prevents the testing of underlying hypotheses (normality, homoscedasticity). Multivariate methods, such as Principal Component Analysis (PCA) or Partial Least Squares-Discriminant Analysis (PLS-DA), often used to analyze metabolic data, do not take into account data’s temporal structure, resulting in a loss of information when used alone. In this study, we applied a method combining ANOVA and PCA: A-SCA (Anova-Simultaneous Component Analysis): SCA is a generalization of PCA taking into account the experimental design, as well as the relationship between variables, to allow data modelisation. Data were first separated into blocks corresponding to the different sources of variation (experimental design factors). Then SCA was independently applied on each block, and permutations test was used to evaluate the significance of model parameters. This method was applied to the study of the effects of low doses of bisphenol A (BPA) on global metabolism in SD rats exposed in the perinatal period, (NIHEs project #5RCE501822). Pregnant rats were exposed to DMSO (vehicle-control), 0.25, 2.5, 25, 250 ng BPA/kg BW/day. Serum samples of the F1 generation were collected on days 21, 50, 90, 140 and 200 of the experiment, and submitted to 1H NMR spectroscopy. Using the A-SCA method, time effects were demonstrated.

551 A Systems Chemical Biology Approach to Predict Effects from Chemical Cocktail Exposure.

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Purpose: In its report, Toxicity Testing in the 21st Century, the National Research Council called for the development of new approaches to human health risk assessment that would rely, in part, on computer-based models rather than animal testing and epidemiology (National Research Council 2007). Using a systems chemical biology approach, we have studied the five commonly used pesticides epoxiconazole, mancozeb, prochloraz, procymidone, and tebuconazole that are all known to have effects on male reproductive development. The purpose of this study was to apply a method previously described by Audouze and Grandjean (Audouze & Grandjean, 2011) to generate hypotheses on the mechanisms linking relevant human outcomes to specific compounds.

Methods: An integrative systems biology approach was adapted to investigate similar or shared modes of action for the chemicals in the mixture of the five pesticides. Human chemical-protein/gene associations were collected from the databases ChemProt, STITCH, and CTD. These associations were then enriched using known protein-protein interactions (PPIs) to yield associations between the chemicals and PPI networks. Subsequently, data integration was performed using in-house tools to extracdata to human diseases and pathways affected by the selected chemicals.

Results: The applied method on these chemicals is capable of identifying known effects on male reproducivity and of predicting novel effects on human health. With the successful identification of known effects, a thorough data analysis is likely to help the linkage between the investigated compounds and their potential effects on other aspects of human health.

Perspectives: As the systems chemical biology approach relies on existing data it serves as an important tool in the efforts to apply alternative methods to animal testing. Furthermore, the method integrates data from many sources, which allows for unravelling of previously unidentified pathways of toxicity.

552 Modeling Species-Specific Metabolic Responses to TCDD in Mice and Rats Using a Reconstructed Metabolic Network.

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2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) elicits species-specific transcriptional and metabolomics responses. Using a reconstructed metabolic network and published microarray datasets, we examined the ability of the network to predict hepatic metabolic responses based on TCDD-elicited differential gene expression profiles. Previously published temporal whole-genome microarray datasets (2-168h) for TCDD-treated female C57BL/6 mice (30 µg/kg) and female Sprague-Dawley rats (rat 200 µg/kg) were used. Context-specific metabolic networks were generated using the Gene Inactivity Moderated by Metabolism and Expression (GIME) method by including induced genes (fold change ≥ 1.5, P1(0) ≥ 0.99) and reactions necessary to maintain a steady-state flux. The mouse context-specific metabolic network revealed more reactions involved in fatty acid (FA) activation and triglyceride synthesis suggesting an increased flux towards lipid synthesis relative to the rat. Meanwhile, mice had fewer reactions involved in FA oxidation, carnitine shuttling, and glycerophospholipid metabolism suggesting FA catabolism is induced in rats exposed to TCDD. Moreover, the predicted induction of glycerophospholipid metabolism is consistent with reported changes in phosphoethanolamine in mice, and phosphocholine and phosphatidylserine in rats. In summary, we show that context-specific metabolic networks based on transcriptomic data are consistent with observed species-specific metabolic changes elicited by TCDD including hepatic lipid accumulation observed in mice and species-specific glycerophospholipid metabolism effects. Moreover, the reconstructed metabolic network facilitated differential gene expression interpretation by providing valuable insights into the species-specific disruption of metabolic responses. Funded by SRF P42E0504911.

553 Multiscale Modeling for Individualized Spatiotemporal Prediction of Drug Effects.


Recently, in silico models for biokinetics have become a relevant way to test the distribution and metabolism of substances in single cells and in organisms. Its relevance is accentuated by the fact that in the next future an animal free toxicity assessment must be implemented in the European Union. Inside the COSMOS project (HEALTH-F5-2010-266835), which is funded by the European Union and a consortium of cosmetic industries (Cosmetics Europe), the central aim is to develop computational methods and in silico models for the prediction of long term toxicity upon repeated exposure to compounds. The power of such methods is (i) to derive quantitative predictions and (ii) to extrapolate results from in vitro measurements into in vivo predictions.

In this contribution, we present a case study that addresses acetaminophen induced hepatotoxicity. Based on a metabolic network model assigned to individual cells, which includes transport, drug metabolism, and intracellular mechanisms leading to the enzymes, we were able to predict the metabolism of acetylaminoeph and production of toxic substances in the liver lobules in different time periods. Depending on the accumulation of these substances, the cell integrity in the liver was estimated. Such estimation is relevant not only for the prediction of physical changes in the organ but also for the prediction of changes in the clearance of the liver under high or repeated drug dosage. Our model also predicted variations of drug toxicity depending on alterations in metabolic enzyme activities. Variations in enzyme activities reflect genetic characteristics or diseases of individuals, thus allowing stratified or even personalized predictions of drug toxicity as well as drug efficacy.
Several polyatomic polycyclic aromatic hydrocarbons (PAHs) are bioavailable to the fetus in utero, can generate DNA damage after transplacental exposure and result in increased incidence of carcinogenesis in offspring in an arylhydrocarbon (AhR)-dependent manner. However, little is known about the effects of environmental PAH mixtures or mechanisms of carcinogenesis. Therefore, we utilized a mouse model of transplacental carcinogenesis to investigate the effect of PAH mixtures collected from environmental samples compared to the known carcinogen, dibenzo[def,p]chrysene (DBC). AhR<sup>−/−</sup> (responsive) females were bred with AhR<sup>−/−</sup> (non-responsive) male mice at eight weeks of age. Females were gavaged on gestation day 17 with DBC, an artificial atmospheric PAH mixture or PAH extract collected from Portland Harbor Superfund Site. Subsamples from each treatment (n=4) were euthanized 24 hrs post-treatment to collect fetal thymus, lung and liver for microarrays. At 3-6 months of age, 79% of the offspring from DBC-treated dams were euthanized due to severe T-cell lymphoma, as expected, which was not detected in the other treatment groups. Exposure of mice to both DBC and the artificial atmospheric mixture resulted in significant (p<0.05) increase in the overall incidence of lung nodules, 100% and 46%, respectively, in offspring at 10 months of age compared to controls. Global transcriptional analysis of fetal tissues 24 hrs post-treatment results in treatment-dependent, tissue-dependent and genotype-dependent gene signatures. A pathway-based approach was utilized to identify mechanisms of exposure and link gene signatures to tumor outcome. Network and transcription factor analysis of the gene clusters further resulted in identification of upstream regulators associated with PAH-induced carcinogenesis. These data describe potential mechanisms for DBC and PAH mixtures in vitro that may be linked to downstream carcinogenesis in offspring. Supported by P42_ES016465.

We contend that functionally conserved toxicity pathways underlie adverse cellular responses to toxicants. Therefore, we are using functional profiling in *Saccharomyces cerevisiae*, to identify the pathways involved in cellular toxicity. In this approach, the ~4500 viable deletion strains are pooled, grown for multiple generations in the presence of a toxicant, and sensitivity of each strain is individually quantified. If a deletion strain is significantly sensitive/resistant, then it provides evidence that the gene product absent in that strain plays a functional role in toxicity. We utilized this approach to identify key susceptibility genes to a wide variety of toxicants. We then applied reverse engineering to infer functional relationships between gene products and assemble functional networks. We used the network inference ARACNE algorithm (using a mutual information threshold of 0.35 (equating to a p-value of 10E-0.25), respectively. Surprisingly, the levels of Cu in cord blood were significantly higher than those in maternal blood with cord/maternal ratios of 1.56 ± 0.13 and 1.41 ± 0.25, respectively. Compared to controls, global transcriptional analysis of fetal tissues 24 hrs post-treatment results in treatment-dependent, tissue-dependent and genotype-dependent gene signatures. A pathway-based approach was utilized to identify mechanisms of exposure and link gene signatures to tumor outcome. Network and transcription factor analysis of the gene clusters further resulted in identification of upstream regulators associated with PAH-induced carcinogenesis. These data describe potential mechanisms for DBC and PAH mixtures in vitro that may be linked to downstream carcinogenesis in offspring. Supported by P42_ES016465.

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Objective: Our previous study indicated that the ambient levels of nickel (Ni), copper (Cu), arsenic (As), and selenium (Se) in Jinchang were 76, 25, 17, and 7 fold higher than Zhangye, respectively. This pilot study was conducted in both cities to assess potential adverse effects of prenatal exposures to these high levels of pollutants on the fetus. Methods: A total of 60 healthy nonsmoking pregnant women were recruited in Jinchang and Zhangye for collection of paired maternal and cord blood samples. ICP-MS was used to measure the concentrations of Ni, Cu, As, Se, and Co in blood samples. ELISA kits were used to measure human C-reactive protein (CRP), interleukin 6 (IL-6), nitrotyrosine, and cotinine. Results: All metals measured were elevated in both maternal and cord blood collected from Jinchang as compared to those from Zhangye, while significant differences were only detected for As, Co, and Se. In addition, Ni and As in cord blood were significantly higher than those in maternal blood with cord/maternal ratios of 1.56 ± 0.13 and 1.41 ± 0.25, respectively. Surprisingly, the levels of Cu in cord blood were significantly lower than paired maternal blood (cord/maternal ratios of 0.41 ± 0.04). The cord blood level of nitrotyrosine in Jinchang was significantly higher than that in Zhangye (p<0.0001). The other inflammatory biomarkers were not significantly different between these two groups. In contrast, CRP in maternal blood was 9 fold higher than cord blood (p<0.0001). The levels of IL-6 and nitrotyrosine were higher in cord blood compared with maternal blood although the differences were not statistically significant. Conclusion: The results suggest that the blood-placenta barrier may effectively prevent transportation of Cu from mother to fetus. On the other hand, Ni and As may easily penetrate placenta barrier and accumulated in cord blood, which may be responsible for the up-regulation of nitrotyrosine.

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A comprehensive survey of chemical, diet and genetic perturbations that activate PPARalpha in the mouse liver has not been carried out but would be useful to identify factors that may contribute to PPARalpha-dependent liver tumours. A gene signature dependent on PPARalpha activation was identified by comparing the transcript profiles after exposure to three PPARalpha activators (Wy-14,643, farnesoid X receptor (FXR) and PPARalpha agonists (1 μM)) in C57BL/6 mice. In independent experiments using transcript profiles from the livers of chemically-exposed male or female mice, the signature correctly predicted activation by 2 known PPARalpha activators but not 10 activators of other pathways. Individual genes in the signature (i.e., Cyp4a10, Fkal) were used in RT-PCR experiments to show the specificity of the response to PPARalpha activators compared to chemical-induced activation of other xenobiotic-activated transcription factors. The signature was used with standard classification methods to identify perturbations in which PPARalpha was activated in an Affymetrix cocompendium of ~750 mouse liver transcript comparisons encompassing a broad range of chemical, dietary and genetic perturbations. We found that PPARalpha is activated by a number of novel chemicals, dietary regimens and genetic mutations. Specific findings include activation by 1) chemicals that cause steatosis (e.g., TCDD, BaP), 2) dietary regimens of triglycerides, “fast food” and “cafeeteria” diets, and fish oil, and 3) nulligrous mutations in a number of genes (Mip2, Erc1, Pot, Hif1α, Fad1/p21, U2D, Hif1α, Hif2αa) that could be secondary to steatosis. The findings increase our understanding of the factors that impact PPARalpha activation and that could contribute to increases in PPARalpha-dependent liver tumours. (This abstract does not represent EPA policy.)

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Nickel (Ni), cadmium (Cd) and chromium (Cr) are human lung carcinogens but the key events leading to their toxic effects are not fully elucidated. In addition, little is known about the molecular basis and the underlying toxic mechanisms of co-exposure of these carcinogenic metals which are frequently observed in environmental exposures. In this study, we investigated the biological responses and effects of the exposures of three metals, individually and in combinations, in BEAS-2B human lung epithelial cells. Utilizing two dimensional gel electrophoresis and mass spectrometry, we identified 455 differentially expressed proteins in BEAS-2B cells exposed to the metals and their mixtures at different doses. The identified protein changes were validated using Western blot and protein functional assays. These altered proteins were mapped to protein interaction networks, toxicity lists, and...
Decabromodiphenyl ether (BDE-209) is a fully brominated diphenyl ether compound used as a flame retardant in polystyrene applications such as casings for computers, upholstery textile and televisions. It is highly lipophilic and persistent and as such it is prone to bioaccumulation and biomagnifications in the food chain. Increasing concentrations of polybrominated diphenyl ethers in wildlife have been documented since the mid 1990s. BDE-209 is a very large molecule, and the general thought was that due to its size it was not bioavailable, and therefore could not bioaccumulate in wildlife. Several studies now have proven this concept wrong, as BDE-209 has been found in many aquatic and terrestrial organisms. Exposure to BDE-209 has been linked to decreased learning and memory in mice, behavioral changes in mice and inhibition of neural stem cells into neurons in culture. As many of the toxic effects of BDE-209 have been found on laboratory exposures, here we analyze the potential effects of an environmentally relevant dose of BDE-209 on zebrafish (Danio rerio) embryo. Zebrafish embryos were exposed to sediment linked to potential developmental and neurotoxic effects on zebrafish embryos.

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Some studies suggest that genotoxic effects of the combustion related aerosol are induced by carcinogenic polycyclic aromatic hydrocarbons (c-PAHs) and their derivatives forming organic fraction of the ambient air particulate matter (PM). The proportion of the organic fraction in PM is known to vary with particle size. The ultrafine fraction is hypothesized to be the most important carrier of c-PAHs, since it possesses the highest specific surface area of PM. To test this hypothesis, the distribution of PAHs in the organic extracts (EOMs) was compared from 4 size fractions of ambient-air aerosols: coarse (1<dₐe<0.5 μm), upper (0.5<dₐe<1 μm), and lower (0.17<dₐe<0.5 μm) accumulation, and ultrafine (dₐe<0.17 μm). High-volume aerosol samples were collected consecutively in 4 localities differing in the extent of environmental pollution. The genotoxicity of EOMs was measured by the analysis of DNA adducts induced in the acellular assay consisting of calf thymus DNA with/without rat liver microsomal 59 fraction coupled with 3P-postlabelling.

The U.S. Air Force is pursuing development of alternative fuels to augment or replace JP-8 jet fuel. Hydroprocessed Esters and Fatty Acids-Mixed Fats (HEFA-F) jet fuel was administered as an aerosol and vapor mixture to 5 male and 5 female Fischer-344 rats per group. Inhalation exposures lasted 6 hours per day for 1 day (with and without an 11-day recovery period), 5 days or 10 days (5 days per week for 2 weeks). Concentrations for each exposure were 0, 200, 700 and 2000 mg/m³; mean aerosol percents were 0, 7, 22 and 28%, respectively. There were no significant changes in male or female body weights due to HEFA-F exposure at any time point. Food and water consumption, urinalysis, clinical pathology, lung cytokine/chemokine and blood cytokines were measured at 10 days. Decreased mean food consumption in male rats was significant at 2000 mg/m³. A 10% decrease in mean water consumption was observed in all HEFA-F exposed female rats. Urinalysis changes included a slight decrease in pH in all exposed rats, as well as a small elevation in ketones and the presence of leukocytes and hemoglobin in the 2000 mg/m³ males. There were no changes in standard clinical chemistry or hematology parameters. Caudal lung tissue was analyzed for cytokines, chemokines and receptors and receptors using a PCR array for rat inflammatory cytokines and receptors kit; no significant changes were seen. Proinflammatory blood cytokines showed no significant differences, although a trend for increasing MCP-1 with increasing dose was seen. Male kidney weight increases were likely related to hyaline droplet formation, relevant only to male rats. Nasal cavity changes included olfactory epithelial degeneration at 2000 mg/m³. Alveolus inflammation was seen in the 2 higher doses at 10 days. To examine sensory irritation, male Swiss-Webster mice were exposed nose-only to 2000 mg/m³ HEFA-F for 30 minutes, resulting in 23% respiratory depression.
Global Gene Expression Changes Induced by Organic Extracts of Air Pollutants in Human Lung Cells.

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Many adverse effects of the ambient air pollution have been linked to polymeric aromatic hydrocarbons (PAHs) occurring in the complex mixture adsorbed onto airborne particles. Besides their genotoxic effect, some of them are known to act via aryl hydrocarbon receptor (AhR)-mediated nongenotoxic and tumor promoting mechanism. This study employed human lung adenocarcinoma cells (A549) to investigate the effect of complex mixture of the air pollutants on the global gene expression changes. We determined whole-genome gene expression profiles (Illumina platform) of cells treated with organic extracts (EOMs) from particulate matter (<2.5 μm) collected in four localities in the Czech Republic differing in the level of the air pollution. Simultaneously, detailed chemical analysis of EOMs from each locality was performed. Despite various sources, EOMs exhibited equal qualitative composition although the absolute level of PAHs and other pollutants in the most polluted locality was much higher comparing to other localities. Gene expression profiles of cells treated with equal amounts of EOM from each locality showed similar patterns. No significant diversity among localities was found. Goeman's global test and KEGG pathway database were applied to identify the most deregulated pathways and contributing genes. We observed significantly deregulated processes and genes common for all localities including metabolism of amino acids such as glycine, serine and threonine (SHMT2, PSAT1), xenobiotic metabolism (CYP1B1, ALDH3A1), vitamins B metabolism (PDXK), TGF-beta signaling (SMAD3), immune system and infectious diseases (IL8, PTGS2) or cell cycle (E2F2, CCND3). It has been proposed that most of them are modulated by activated Ah-receptor. Our results suggest the prominent role of activated Ah-receptor and other events possibly leading to the metabolic reprogramming and tumour promotion in A549 cells. Support: Grant Agency of the Czech Republic (CZ: P503/11/0142).

Evaluation of Additivity of Binary Mixtures of Perfluoralkyl Acids (PFAAs) on Peroxisome Proliferator-Activated Receptor-Alpha (PPARα) Activation In Vitro.

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Perfluoralkyl acids (PFAAs) are found globally in the environment and in animal tissues, and are present as mixtures of PFAA congeners. Mechanistic studies have found that in vivo effects of PFAAs are mediated by PPARα. Our previous studies showed that individual PFAAs activate PPARα transffected into COS-1 cells. Here we evaluated whether binary combinations of perfluorooctanoic acid (PFOA, C8) and other PFAAs interact in an additive fashion to activate PPARα. COS-1 cells in 96 well plates were transiently transfected with mouse PPARα luciferase reporter plasmid. After 24 hours, cells were exposed to either vehicle control (0.1 % DMSO or water), PPARα agonist (WY14643, 10 μM), C8 or perfluorononanoic acid (C9) at 1 - 128 μM, perfluorohexanoic acid (C6) at 8 – 1024 μM, or perfluorooctane sulfonate (C8S) at 4 – 384 μM to generate sigmoidal dose-response curves. In addition, cells in the same plate were exposed to binary combinations of C8 + either C6, C9, or C8S, in an 8x8 factorial design. After 24 hours of exposure, cells were lysed and luciferase activity was measured. Data were transformed on a fold-induction and % maximal response basis. The dose-response data for individual chemicals were fit to sigmoidal curves and analyzed with nonlinear regression to generate EC50s and Hill slopes, which were used in response-addition and dose-addition models to calculate predicted responses for mixtures. All PFOA+PFAA combinations produced dose-response curves that were closely aligned with the predicted curves for both response addition and dose addition. However, at higher concentrations of all chemicals, the observed response curves deviated upward from the predicted models of additivity, although with more variability. We conclude that at the lower concentration ranges, binary combinations of PFAAs behave additively in activating PPARα in the COS-1 cell system. This abstract does not necessarily reflect USEPA policy.

Reference Dose (RID)-Based Chronic Human Health Hazard Ranking System for Complex Mixtures—Assessment of Polar Nonhydrocarbons in Groundwater at Biodegrading Petroleum Sites.

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Chronic human health hazard evaluations of chemical mixtures are challenging and generally rely on similarities in toxicity mechanisms and dose-additivity of chemicals in the mixture. However, such assessments are inadequate for highly complex mixtures such as those present in groundwater at biodegrading petroleum sites where thousands of compounds representing many distinct structural classes of chemicals may be present. A hazard ranking system based on USEPA Reference Doses (RIDs) was developed to evaluate the potential chronic human health hazards presented by complex mixtures of polar petroleum biodegradation products in water. Equivalent risk-based drinking water concentrations for the range of identified RIDs were derived using the USEPA Regional Screening Level (RSL) equation for Tap Water Screening Levels (SLwater-nc-ing). Based on the RIDs of representative chemicals for each structural class of potential biodegradation products, overall summary hazard rankings of “Low”, “Low to Moderate”, and “Moderate”, were assigned. Classes constituting chemicals with RIDs ≥ 0.1 mg/kg/day were defined as being of “Low” hazard; “Low to Moderate” if 0.1<RID≤ 0.01 mg/kg/day, and “Moderate” if 0.01<RID< 0.001 mg/kg/day. These three groups included essentially all of the potential polar biodegradation products for which RIDs were available. This RID based system is consistent with similar systems developed by USEPA OPPTS and UN GHS programs and was validated by review of USEPA summary documents.

The ranking system was applied to groundwater samples collected from biodegrading petroleum sites and the results show that the vast majority of the polar biodegradation products are in structural classes that may present “Low” hazard to humans. Overall, the polar mixtures are unlikely to present a significant risk to human health if consumed as drinking water.
and N.

Monroe, LA; 3Pharmacology, Toxicology & Neurosciences, LSU Health spectrums of the crude oils obtained with EPR and 1H- and 13C-NMR. Sources of oil oil from several sources and correlate these with peaks of select constituents from Here, we identify several toxicological effects upon acute exposure of rats to crude cyclic aromatics, nitrogen, sulfur, asphaltene and porphyrin nickel and vanadium. Crude oil varies with source in its relative composition of alkanes, simple and poly- Crude oil varies with source in its relative composition of alkanes, simple and poly- radic and vanadium porphyrin and lowest in benzene, most affected liver weight lesser effect on spleen and liver weights and ALP. Iraqi oil, richest in asphaltene free peak with intensity greatest for Iraqi oil. Vanadium porphyrin was detected in Iraqi peroxisomal beta-oxidation and NADPH oxidation pathways. Our results indicate that Iraqi crude oil has a potential to induce liver in oxidative stress and DNA damage in rats.

PS 568 Different Toxicity Outcomes in Rats Correlated with EPR and NMR Spectra of Crude Oil from Various Sources.

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Crude oil varies with source in its relative composition of alkanes, simple and poly- cyclic aromatics, nitrogen, sulfur, asphaltene and porphyrin nickel and vanadium. Here, we identify several toxicological effects upon acute exposure of rats to crude oil from several sources and correlate these with peaks of select constituents from spectra of the crude oils obtained with EPR and 1H- and 13C-NMR. Sources of oil were Louisiana sweet crude, Nigerian (Qua Iboe) sweet crude and Iraq high sulfur (ONTA, Inc., Toronto, Canada). Female Sprague-Dawley rats were given 2 daily doses of 2.5 and 5 ml/kg of oil or vehicle (0.5% DMSO in corn oil) by oral gavage. Rats were euthanized after 48 h and blood was taken for hematology, clinical chem- istry and immunoassay for cytokines and vasoactive agents. Femur bone marrow cells were assayed for CFU-GM myeloid progenitors. All oils elevated serum alka- line phosphatase (ALP) with LA oil being least active, in agreement with liver pathology. Liver weight increased 25-75% and CYP1A1 protein was elevated with all, Iraqi oil being most effective. Granulocytes increased -2-fold by high dose of LA and Iraqi oil. Spleen weights decreased 30% with high dose of Iraqi and Nigerian oil. CFU-GMs decreased only with high dose of LA oil (40%). Vasocostructor endothelin-1 increased -2-fold in serum from rats treated with Iraqi and Nigerian oils. 1H- and 13C-NMR spectra gave qualitatively similar alkane peaks for all oils, but differing aromatic peaks, e.g., benzene (1H-NMR 7.38 ppm) of Iraqi oil was least intense. EPR spectra of all oils exhibited asphaltene free radical peak with intensity greatest for Iraqi oil. Vanadium porphyrin was detected in Iraqi oil only. Summarizing, LA sweet crude oil was uniquely myelosuppressive and with lesser effect on spleen and liver weights and ALP. Iraqi oil, richest in asphaltene free radical and vanadium porphyrin and lowest in benzene, most affected liver weight and CYP1A1 (LA Board of Regents).

PS 569 Complex Mixtures of PAHs Are More Potent Than Benzo[a]pyrene at Inducing Cellular Inflammatory and DNA-Damage Response.

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Complex mixtures of polycyclic aromatic hydrocarbons (PAHs) are present in air particulate matter (PM) and have been associated with many adverse health effects including cancer and cardiovascular disease. We hypothesize that interactions between different PAHs stand for a major biological effect. We have previously showed that soil PAH extracts induce persistent DNA damage including prolonged activation of DNA damage markers H2AX and Chk1. To further study the effects of mixtures of PAHs biological testing of air PM extracts were performed using human HepG2 and H1395 cells and sensitive cellular endpoints relevant for car- cinogenesis (DNA damage response) and cardiovascular pathogenesis (inflamma- tory signaling). The cellular response was compared to benzo[a]pyrene (BP) and dibenzo[a,l]pyrene (DBP). The results showed a more than additive response for binary mixtures of BP and DBP on activation of Chk1. Persistent activation of DNA damage signaling was observed at significantly lower concentrations of air PM ex- tracts than BP alone. Activation of DNA damage signaling was more persistent in air PM fractions containing PAHs with more than four aromatic rings suggesting larger PAHs contributes to the genotoxicity of PAHs in air PM. Furthermore, we observed significant up-regulation of pro-inflammatory stress responsive genes including several cytokines in response to air PM extracts corresponding to 1 nM BP. Employing specific inhibitors showed that the activation of inflammatory sig- naling was mediated through the Ah-receptor and MAPK signaling. Taken to- gether, our data indicate synergistic events due to PAH interactions. This suggests that human health risk assessment based on additivity such as toxicity equivalency factor scales may significantly underestimate the risk of exposure to complex mixtures of PAHs.

Perfluoroalkylated and polyfluoroalkylated substances (PFASs) are a large class of chemicals that have emerged as global environmental contaminants. This study, carried out in accordance with the European chemicals legislation (REACH) guidelines for risk assessment, evaluated possible health risks of 17 PFAS congeners in the Swedish population. The exposure assessment was based on blood and serum levels from biomonitoring studies in the Swedish population. Population groups consid- ered were the general population and occupationally exposed professional ski- ers. The hazard assessment primarily considered hepatotoxicity and reproductive toxicity which were endpoints shared by the selected congeners. Read-across was performed to the closest most potent congener for 12/17 congeners lacking toxico- logical data and/or internal dose levels. The result of the risk characterization showed no cause for concern for hepatotoxicity or reproductive toxicity in the gen- eral population, except for hepatotoxicity in a subpopulation eating PFOS-contam- inated fish. However, a cause for concern was identified for the non-conventional endpoints disrupted mammary gland development and immunotoxicity. For the occupationally exposed professional ski waxes safe use could not be shown based on concern for liver toxicity by PFOA and by all congeners in combination, for re- productive toxicity by all congeners in combination as well as for disrupted mam-mary gland development and immunotoxicity. This is the first attempt to assess the health risks to a combination of a large number of PFASs.

PS 570 Individual and Cumulative Health Risk Assessment of 17 PFASs in the Swedish Population.

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Genotoxic carcinogens are present in the human diet, and Benzo(a)pyrene (BaP), 2-Amino-1-ethyl-6-phenylimidazo[4,5-b]pyridine (PhIP) and acrylamide (AC) represent three important examples. BaP is a polycyclic aromatic hydrocarbon generated by incomplete combustion of organic substances such as lipids, thus con- taminating numerous foodstuffs. PhIP is a heterocyclic amine formed when meat is cooked, and AC forms when foods, such as potatoes and cereals, are cooked at temperatures exceeding 100°C. Individually these have been shown to be genotoxic but the biological consequences of exposure to mixtures of these chemicals have not been systematically examined.

The aim of the current study was to examine the biological response of MCL-5 cells (metabolically competent human lymphoblastoid cell line) to mixtures of these genotoxins at concentrations relevant to human exposure (μM to sub-nM). Cells were exposed to the chemicals individually or in mixtures for 24h and mutagenicity was assessed through resistance to trifluorothymidine at the thymidine kinase (TK) locus and 6-thioguanine at the hypoxanthine-guanine phosphoribosyltransferase (HPRT) locus.

At the TK locus the mixtures produced non-monotonic mutation responses; 100nM BaP combined with some low, non-mutagenic concentrations of other mutagens showed synergism, while antagonism was observed for 10μM BaP in mix- tures with higher concentrations of other genotoxins. Responses differed between the two loci, with a higher than anticipated mutation frequency (MF) observed for high concentration combinations at HPRT compared to TK. Combining 10μM BaP with 100nM PhIP reduced the number of cells in S phase with a correspon- ding increase in sub-G1 and G1. Moreover, ethoxyresorufin-O-deethylase (EROD; CYP1A) activity and CYP1A1 mRNA levels significantly correlated with each other and with the MF at TK emphasising the involvement of the CYP1A family in this mutation response. This non-monotonic MF is of significance when consider- ing risk assessment, especially at low concentration combinations where the indi- vidual chemicals are not measurably mutagenic.

R. David and N. J. Gooderham. Imperial College, London, United Kingdom.

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571 Mixtures of Benzo(A)Pyrene with Direct or Indirect Acting Mutagens Have a Nonmonotonic Mutation Profile.

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At the TK locus the mixtures produced non-monotonic mutation responses; 100nM BaP combined with some low, non-mutagenic concentrations of other mu-
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When assessing risks posed by environmental chemical mixtures, whole mixture approaches are preferred to component approaches. When toxicological data on whole mixtures as they occur in the environment are not available, EPA guidance states that toxicity data from one or more mixtures considered “similarly sufficient” to the environmental mixture(s) can serve as surrogates. However, the selection process of which mixtures to experimentally evaluate is an open line of inquiry. The objective of our study was to demonstrate a proof-of-concept strategy for the selection process using a mixture of 17 polychlorinated dibenzo-p-dioxins (PCDDs), polychlorinated dibenzofurans (PCDFs) and polychlorinated biphenyls (PCBs) based on (i) human biomonitoring data from the National Health and Nutrition Examination Survey (NHANES); and (ii) single chemical dose response models from the literature. Principal component (PC) analysis was conducted to identify patterns of chemicals present in humans as measured in serum concentrations in the US population using NHANES data. Five different concentration patterns were found to represent 82% of the variation in the mixture concentrations for these 17 chemicals. Population-based median concentration estimates for the chemicals identified by large absolute values in the PCs were used to define mixing proportions for the five mixtures. Robust optimal designs, using a maximin optimization criterion which is robust to model misspecification, were determined for the mixtures using published dose-response models in Crotfor et al (2005, EHP). The approach provides design information for conducting mixtures studies in support of a whole mixtures risk assessment. Next steps include estimating the dose values required to approximate the blood concentrations in the mixtures. Determining how representative these mixtures are to the population concentrations while testing for similar sensitivity is a long-range research goal. (The authors gratefully acknowledge the support from #R01ES015276, #T32 ES007334, and #U1TR000558.)

573 Coexposure to Low-Dose Model Testicular Toxicants Induces Gene Alterations.
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Testicular effects of chemical mixtures may differ from those of the individual chemical constituents. This study assesses the co-exposure effects of the model germ cell- and Sertoli cell-specific toxicants, X-radiation (x-ray) and 2,5-hexadienoic acid (HD), respectively. X-ray induces germ cell apoptosis and HD has been shown to attenuate the x-ray effect on germ cells. Adult rats were exposed to different levels of x-ray (0.5Gy, 1Gy, 2Gy), HD in the drinking water for 18 days (0.33%, 1%, alone or in combination. Custom PCR arrays were generated based on a panel of genes identified through microarray studies; hierarchical clustering of these PCR arrays identified dose-dependent treatment effects on the apoptosis-related genes Fas, Aen, and Casp3. The 5Gy + 1% HD co-exposure induced Aen and Casp3 expression, with a maximum fold induction of 3.1 and 2.6, respectively. Also, Fas was significantly induced 2.6-fold at this co-exposure dose, despite no induction of Casp3, Fas, or Aen expression by x-ray or HD alone. In order to assess cell type-specific attenuation of HD effects with x-ray co-exposure, we used laser microdissection (LCM) to examine a panel of apoptosis-related transcripts using PCR arrays. Several pro-apoptotic genes were identified, which increase with a fold change greater than 1.4 after 2Gy x-ray exposure, including Tra2, Tinbl10 and Fas. We also identified an anti-apoptotic gene, Bcl2a1d, which increases in fold change across the different exposures (0.33% HD, 1.3 fold; 2Gy x-ray, 1.6 fold; 0.33% HD + 2Gy, 3.1 fold). Ingenuity Pathway Analysis of the gene expression data produced two over-represented pathways across all treatment groups examined with LCM: Induction of Apoptosis by HIV1 and Death Receptor Signaling. We amplified LCM RNA and examined the same apoptosis transcripts. When compared to our unamplified RNA, we found a Pearson correlation coefficient of 0.64, indicating that use of amplified LCM RNA may introduce significant bias. These results provide insight into environmentally relevant low-dose co-exposures of model testicular toxicants.

574 Development of a Portable In Vitro System for Lab and Field Aerosol Exposure Studies.
J. Zava, K. Lichtveld, S. Ebersviller, G. W. Walters, H. E. Jeffries, K. G. Sexton, W. Vizuet, T. Rusyn and J. Jaspers, Environmental Sciences and Engineering, University of North Carolina at Chapel Hill, Chapel Hill, NC.

There is a growing interest in studying the toxicity of multi-pollutant mixtures found in ambient air, and the U.S. Environmental Protection Agency (EPA) is moving towards setting standards for these types of mixtures. Conventional in vitro exposure methods do not properly emulate human exposures and are not adequate to meet the EPA strategic plan to demonstrate a direct link between air quality and health effects. Exposure of cells at the air-liquid interface (ALI) is the most realistic approach to emulate in vivo exposures. A new portable in vitro system that uses electrostatics to deposit particles onto the cells has been developed and this study demonstrates the efficacy of this device. The portable system provides a viable environment within its microfluidic fields for determining the effects of human lung cell growth on permeable membrane supports. This method provides an efficient and effective way to expose cells to particles without prior collection and subsequent resuspension in a liquid medium. Cell viability testing included maintaining normal cell culture conditions for exposures across the ALI for periods of up to 4 hours outside of a tissue culture incubator. Results from all cell viability testing with A549 human lung cells show that no parts, components, materials or operating characteristics of the portable sampler induce any cytotoxicity, as measured by lactate dehydrogenase (LDH), or inflammatory response, as measured by interleukin-8 (IL-8). A549 cells were then exposed to photochemically-aged diesel exhaust (DE) using the UNC rooftop smog chamber. An exposure dose of 3.6 μg of DE particles in air was delivered to the cells and a 3-fold increase in LDH and IL-8 was observed when compared to controls. This new device can serve as a stepping-stone for researchers and engineers to develop new and improved in vitro exposure technology suitable for field use.

575 Chemical Dispersants Used in the Gulf of Mexico Are Cytotoxic to Human Lung and Skin Fibroblasts.
J. Wise1,2, S. Wise1,2,3, H. Xin1,2,3, J. Griffin4 and J. Wies1,2,3, ‘Wies Laboratory of Environmental and Genetic Toxicology, University of Southern Maine, Portland, ME; 4Maine Center for Toxicology and Environmental Health, University of Southern Maine, Portland, ME; Department of Applied Medical Science, University of Southern Maine, Portland, ME; Department of Coastal Sciences, University of Southern Mississippi, Ocean Springs, MS.

Chemical dispersants are chemical compounds that can be used to aid in the cleanup of crude oil spills. In 2010, they became a significant public health concern due to the BP Deepwater Horizon Oil Crisis; when millions of gallons of chemical dispersants specifically Corexit® 9527 and 9500 were applied via aerial spray and deepwater injection to break up the crude oil. Toxicity of Corexit® to humans is unknown, and the primary routes of exposure to these chemical dispersants are inhalation, direct dermal contact and ingestion. The objective of this study is to determine the cytotoxicity and genotoxicity of these two dispersants in human lung (WTHBF-6) and skin (BjH(TERT) fibroblasts. Cells were treated with and without 59 fractions with cofactors, because fibroblast cells may not readily express cytochrome P450 enzymes necessary to metabolize chemicals. Corexit® 9500 was cytotoxic to lung and skin cells. Specifically in skin, 50, 250, 350 and 500 ppm, it induced 95, 89, 52 and 3 percent relative survival, respectively. 59-mediated metabolism increased toxicity inducing 78, 84, 39 and 2 percent relative survival, respectively. Corexit® 9527 and 9500 are used to break up crude oil in skin, 500, 650, 850 and 1000 ppm, it induced 89, 56, 73, and 24 percent relative survival, respectively. 59-mediated metabolism increased toxicity inducing 65, 60, 22 and 0 percent relative survival, respectively. Our genotoxicity data is less clear. Corexit® 9527 induced 8 aberrations per 100 metaphases at 850 ppm, but caused cell cycle arrest at 875 ppm. Similarly Corexit® 9500 induced 14 aberrations per 100 metaphases at 500 ppm, but caused cell cycle arrest at 425 ppm. Ongoing and future work will consider the effects of dispersed oil.

576 Micronanoparticle Suspension Formulation Development for Medical Chemical Countermeasures.
L. Cabell1, J. McDonough1, A. Clark1, T. Belkis2 and L. Mobley3, ‘SWRI, San Antonio, TX; ‘CBMS /MITS, Frederick, MD. Sponsor: L. Johnson.

Current U.S. medical chemical countermeasures require battlefield stability and an effective shelf life. One example is bis-pyridinium oximes that are used for the treatment of Organophosphate (OP) intoxication to counter the effects of AChE inhibition. This class of oximes presents poor thermal stability in an aqueous formulation due to hydrolytic cleavage. CBMS and SWRI have developed non-Newtonian suspension formulations of oximes in cottontose oil (CSO) vehicles that impart oxime solid state stability characteristics to oxime/CSO liquid suspension formulations. MM64 DMS and HI-6 DMS have shown superior thermal stability in these liquid suspension formulations compared to aqueous formulations and equal thermal stability compared to solid formulations. These types of formulations have shown bioavailability comparable to parent active pharmaceutical ingredients (API) in aqueous formulations. Cmax and Tmax for these APIs are relatively the same with absorption and elimination not substantially affected. In principle, these types of
suspension formulations can be used to stabilize any drug that is prone to hydrolytic cleavage. In addition, the nature of these suspensions inhibits drug-drug interaction in the formulations as well as promotes differential drug controlled release if designed appropriately. We have designed these suspensions with high zero-shear viscosity, which allows it to resist sedimentation and non-Newtonian behavior that causes shear thinning, such that the viscosity decreases and the product flows easily under a shear force, such as injection through a needle. Furthermore, the viscosity and sedimentation behavior are controlled by formulation parameters such as particle size and concentration. DISTRIBUTION STATEMENT A. Approved for public release; distribution is unlimited.

577 Application of Generalized Concentration Addition to Receptors That Dimerize.
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Concentration addition is a standard toxicological method for analyzing mixtures of similarly acting compounds. A significant limitation of concentration addition is that it cannot be used to describe the effect of mixtures containing partial agonists at response levels above that for the compound with lowest efficacy. Unfortunately, partial agonism is a common phenomenon, limiting the application of concentration addition. We previously proposed generalized concentration addition (GCA) as a solution to this issue and successfully used this method to describe data from experiments involving mixtures of full and partial agonists of the AhR receptor. GCA requires the use of dose-response functions that are invertible over the relevant range. This posed no problem for the AhR-ligand systems described by a Hill function with a Hill coefficient of one. Receptors that dimerize (or receptors that bind two ligands) are often modeled using Hill functions with Hill coefficients of two, but these functions are not appropriately invertible. We used mathematical models based on equilibrium kinetics and mass balance to describe a simple pharmacodynamic model of binding of ligands to receptors followed by dimerization: e.g., ARAR, AR, ARAR, ARRA, where A is a ligand, R is a receptor, AR is a ligand-receptor complex, and ARRA is the dimerized complex. This leads to theoretical dose-response functions that are invertible and different from Hill functions but approximate Hill functions at low doses. Such models may be applicable to examine mixtures of full and partial agonists of several systems of receptors that dimerize including the androgen and estrogen receptors.

578 Generalized Concentration Addition Models Some but Not All Interactions of Mixtures of Androgen- and Estrogen-Receptor Active Compounds in High-Throughput Screening Assays.
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As part of the U.S. Tox21 collaboration, 69 mixtures of compounds previously found to be active in estrogen or androgen receptor (ER or AR) reporter assays were tested in ER and AR high throughput screening assays in agonist and antagonist mode. ER-active compounds used were acetochlor, alachlor, bisphenol A, butyl benzyl phthalate, chlorodecone, DDT, dicumyl peroxide, ethyleneamidine, p-nonylphenol, and zearalenone. AR-active compounds were androstenedione, dexamethasone, fluoxymesterone, hydroxylutamide, medroxyprogesterone acetate, mestranol, oxymetholone, and progesterone. Of the 69 mixtures, 32 were AR-active compounds only, 23 were ER-active compounds only, and 14 were both groups of compounds. Mixtures contained between 2 and 18 chemicals and were equipotent mixtures, mixtures of equal concentration and mixtures that varied potency of the individual chemicals. The 18 compounds were also tested individually. A generalized concentration addition model for partial agonism and antagonism was used to model the concentration-response of the mixtures as a function of the parameters of the response for the mixture components. The parameters were derived using two optimization methods. In the first method, the parameters for the mixtures components were derived by fitting the model to the single-component data, and then the model was used to predict the mixture effects. In the second method, the parameters were obtained by fitting the model to all the data (both single-component and mixture) simultaneously. The second method generally accurately models both the single-component and mixture data, although there were several mixtures that were not accurately represented by the model. The first method was inaccurate for many more mixtures. The experiment demonstrates that high-throughput assays can be used to evaluate dose addition theory for large numbers of individual chemicals and mixing ratios.

579 Cumulative Risk Assessment of Radon and Confounder Mixtures.
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Cumulative risk assessment is a challenging proposition because it involves integration and collective distillation of all available data capturing the essence of the potential risks of multiple stressors, chemical and nonchemical. The conclusions of such assessments must also integrate the inherent disparities, the constraints of the data gaps, and address data uncertainties. More recently communities and stakeholders have also voiced concerns about health effects beyond the single stressor-specific critical effects and the role of interactions in the overall toxicity of real life exposures. Most often the focus has been on chemical toxicity; here we present a case of chemical plus radiological effects. The Agency for Toxic Substances and Disease Registry (ATSDR) has updated the Toxicological Profile for Radon to comprehensively evaluate its systemic toxicity and carcinogenicity. It features information from individual and pooled epidemiological studies of miner and residential cohorts to assess radon carcinogenicity. These studies largely accounted for smoking and arsenic as confounders. Crystalline silica and diesel exhaust carcinogenicity was not integrated into the miner assessments. Thus, radon carcinogenicity may have been overestimated in miners. Residential epidemiological studies of radon have assessed cancer risk addressing smoking as a confounder and suggested that lung cancer mortality in radon-exposed populations may be 25 times higher among smokers than never-smokers. It is not possible to avoid exposure to radon and its progeny since they are ubiquitous. For this reason, to limit potential health effects, EPA recommends keeping radon air level <4 pCi/L in residential settings. Diesel exhaust and other indoor chemicals/non-chemical stressors, including biologics, might enhance the health effects of residential radon and should be considered as part of any risk assessment. (The views expressed in this abstract are those of the authors and do not necessarily represent the views or policies of ATSDR.)

580 90-Day Inhalation Toxicity Study with Hydroprocessed Esters and Fatty Acids Jet Fuel from Camelina (HEFA-C) in Rats.
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The Department of Defense is actively pursuing the development of alternative fuels to augment or replace petroleum-based jet fuels. Towards these efforts, numerous synthetic and biologically-based fuels are currently under consideration for military use. Given the widespread use of fuels within the military, the health effects associated with occupational exposure to fuels remains a concern. Toxicity studies are being performed with the fuels in order to assess any possible health effects related to fuel exposure. Hydroprocessed Esters and Fatty Acids jet fuel (HEFA) is a type of hydrotreated renewable jet fuel currently under consideration. One specific type of HEFA is generated from oils extracted from the camelina plant (Camelina sativa; HEFA-C). In order to evaluate potential toxicity of HEFA-C, an in vivo 90-day whole body inhalation study was performed with the fuel (concentrations of 0, 200, 700 and 2000 mg/m³ for 6h/day, 5 days/week) using male and female Fischer 344 rats. Following exposure, a series of toxicity endpoints was evaluated, including food consumption and body weight, genotoxicity by micronucleus (MN) test, neurotoxicity and histopathology. There was no change in food consumption attributed to fuel exposure and the average body weight was found to slightly decrease (not statistically significant) in animals exposed to the high concentration. MN test was negative for evidence of genotoxicity. No significant effects were observed on clinical chemistry or hematology evaluations and no significant neurobehavioral effects were observed based on functional observational battery and motor activity tests. Minimal effects attributable to exposure to HEFA-C were observed with histopathology. These effects included goblet cell hyperplasia and olfactory epithelium degeneration at the highest concentration of exposure. These data will help guide the establishment of an occupational exposure limit for this alternative jet fuel.
Atrazine is one of the most commonly applied herbicides in the United States. To control broadleaf and grassy weeds, atrazine is used on many agronomic crops. The present study aimed to evaluate the cytotoxic effects of atrazine in HepG2 cells by inhibiting cell proliferation and/or inducing cell death and disrupting mitochondrial membrane potential at concentrations of 50 μM and higher.

Further studies will be performed to better understand the effects of atrazine on HepG2 cells.
585 Effect of p, p'-DDE on Adipogenesis and Macrophage Activation.


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There is evidence that exposure to organochlorine compounds, such as DDE, may contribute to dyslipidemia, obesity, and insulin resistance, a cluster of signs that comprise the metabolic syndrome. Obesity results from adipocyte hyperplasia and hypertrophy which leads to recruitment of immune cells to adipose tissue and eventually contributes to localized chronic low-grade inflammation and adipocyte dysfunction. This study investigated a potential COX-2 dependent mechanism by which DDE could modulate normal physiological function and facilitate an increased risk of developing metabolic syndrome following exposure. Murine (J774.1) and human (THP-1) macrophage cell lines and murine preadipocytes (3T3-L1) were exposed to DDE to determine the effects of DDE on inflammation and adipogenesis. Macrophage cell lines were exposed to DDE or a known COX inhibitor for 18 hr before treatment with an inflammatory challenge of lipopolysaccharide (LPS) or palmitic acid (PA). Cell culture supernatants were analyzed for the presence of prostaglandins PGE2, PGD2, PGF2α, arachidonic acid (AA), and thromboxane (TXB2) and for proinflammatory cytokines, including tumor necrosis factor (TNFα), IL-6, and IL-1β. Preliminary experiments indicated that α,β-unsaturated fatty acids are activated by the cross-linking of FcεRI with or without TNFα, a known COX inducer and suppressor of adipogenesis. Mature adipocytes were analyzed for intracellular lipid accumulation by Oil Red O staining. Both DDE and COX inhibitors increased adipogenesis in a dose-dependent manner in all treatments. In macrophages, DDE exposure followed by LPS or PA challenge reduced secretion PGE2, PGD2, and PGF2α, and caused a slight increase in secretion of TNFα over vehicle. COX enzymes have a complex role in both inflammation and adipogenesis, and these results suggest that DDE exposure may contribute to altered adipogenesis and increased inflammation, potentially through a COX-2 dependent mechanism.

586 Detection of Dichlorvos Targets in Hepatocytes.

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Toxicity of dichlorvos (DDVP), an organophosphate (OP) pesticide, classically results from modification of the sitee in the active sites of cholinesterases. However, DDVP also forms adducts on unrelated targets such as transferin and albumin, suggesting that DDVP causes cellular perturbation by modifying non-cholinesterase targets. Here, we identify novel DDVP target proteins of a human hepatocyte-like cell line (HepaRG) in competitive pull-down experiments using DDVP and a biotin-linked organophosphorus compound (10-fluoroethoxyphosphinyl-N-biotinamidopentyldecanamide; FP-biotin), which competes with DDVP for similar binding sites. Using the competition assay and mass spectrometry, we show that DDVP forms adducts to six new target proteins, including Glyceraldehyde-3-Phosphate Dehydrogenase (GAPDH). We validated the results using purified GAPDH incubated with DDVP and found that GAPDH Tyr-314, a residue in the known active site, is modified by DDVP. DDVP activity is inhibited in a concentration dependent manner in DDVP-treated HepaRG cells, suggesting that DDVP treatment directly inhibits the enzyme. These results may help explain the chronic metabolic effects of DDVP exposure.

Opinions, interpretations, conclusions, and recommendations are those of the authors and are not necessarily endorsed by the U.S. Army. Citations of commercial organizations or trade names in this report do not constitute an official Department of the Army endorsement or approval of the products or services of these organizations. The research described herein was sponsored by the USAMRMC and Military Operational Medicine Research Program. This research was supported in part by an appointment to the Postgraduate Research Participation Program at the U.S. Army Center for Environmental Health Research (USACEHR) administered by the Oak Ridge Institute for Science and Education through an interagency agreement between the U.S. Department of Energy and USACEHR.

587 Organophosphorus Pesticides Activate Mast Cells Ex Vivo and In Vivo.

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Mast cells are immune cells involved in diverse inflammatory diseases and are best known for their critical role in allergic reactions and asthma. In asthma, mast cells are activated by the cross-linking of FcεRI (the high affinity IgE receptor) with IgE-bound allergens. We hypothesized that organophosphorus pesticides (OPs) can activate mast cells in an IgE-independent manner. OPs are known to modulate immune function at levels that do not cause significant inhibition of cholinesterases, and previous studies of total peritoneal cells exposed to the OP malathion at levels that do not cause significant inhibition of cholinesterases, and previous studies of total peritoneal cells exposed to the OP malathion at levels that do not cause significant inhibition of cholinesterases, and previous studies of total peritoneal cells exposed to the OP malathion at levels that do not cause significant inhibition of cholinesterases, and previous studies of total peritoneal cells exposed to the OP malathion at levels that do not cause significant inhibition of cholinesterases, and previous studies of total peritoneal cells exposed to the OP malathion at levels that do not cause significant inhibition of cholinesterases, and previous studies of total peritoneal cells exposed to the OP malathion at levels that do not cause significant inhibition of cholinesterases, and previous studies of total peritoneal cells exposed to the OP malathion at levels that do not cause significant inhibition of cholinesterases.

In vivo treatment with paraoxon induced β-hexosaminidase, a marker for mast cell degranulation, in percoll gradient-purified mast cells, suggesting that the modification directly inhibits the enzyme. These results may contribute to altered adipogenesis and increased inflammation, potentially through a COX-2 dependent mechanism.

Thiamethoxam (TMX), one of the most important insecticides, is carcinogenic and hepatotoxic in mice but not in rats. Green et al. (Toxicol. Sci. 86, 36-47, 2005) established that TMX is a much better substrate for mouse liver cytochrome P450s than the corresponding rat or human enzymes in forming desmethyl-TMX (dm-TMX), which is also hepatotoxic, and clothianidin (CLO), which is not hepatotoxic or carcinogenic. The present investigation examines the hypothesis that liberation of formaldehyde (HCHO) or an N-methylol intermediate during TMX and dm-TMX metabolism to CLO and dm-CLO, respectively, provides a molecular explanation for the mouse-specific toxicity. Mouse liver microsomes were compared to rat and human liver microsomes in the metabolism of the seven commercial neonicotinoids and dm-TMX in the presence or absence of the cofactor, NADPH. HCHO production was determined by derivatization with 2,4-dinitrophenylhydrazine and monitoring by HPLC while other TMX metabolites (dm-TMX, CLO and dm-CLO) were quantitated by LC/MS. TMX and dm-TMX yielded more HCHO than any of the six other commercial neonicotinoids and mouse microsomes gave much higher conversion than rat or human microsomes. HCHO or an N-methylol intermediate in TMX and dm-TMX metabolism is therefore the candidate reactive hepatotoxicant. The N-methylol intermediates are difficult to analyze because of their chemical instability. These results provide HCHO liberation as an alternative to the earlier explanation of TMX hepatotoxicity in which dm-TMX is the proposed reactive metabolite and its toxicity is exacerbated by dm-CLO inhibition of inducible nitric oxide synthase (iNOS) (Green et al., 2005). Although the HCHO generation proposal versus the dm-TMX/iNOS inhibition hypothesis for TMX toxicity is not fully resolved, the mouse vs. rat vs. human comparison of TMX metabolism reconfirms that the rat (showing no carcinogenic effects) is the best risk assessment model for humans.

589 Neonicotinoid Insecticides: Formaldehyde Generation As A Proposed Mechanism of Mouse-Specific Hepatotoxicity and Hepatocarcinogenicity of Thiamethoxam.

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Chiral Pesticide Pharmacokinetics: A Range of Values.


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Approximately 30% of pesticides are chiral and used as mixtures of two or more stereoisomers. In biological systems, these stereoisomers can exhibit significantly different pharmacokinetics (absorption, distribution, metabolism, and elimination). In spite of these differences, these "mixtures" are often treated as a single compound in exposure and hazard assessment. To evaluate the impact of pharmacokinetic differences on internal exposure (dose) estimates, we studied the stereoisomer-specific metabolism of twenty chiral 1,2,4-triazole fungicides in hepatic microsomes, hepatocytes, and purified cytochrome P450s (P450). Additionally, we measured the metabolism of the pure stereoisomers for three of the
fungicides. Studies indicated that CYP3A4 was a major metabolizing enzyme for a majority of the fungicides. In general, the stereoisomers of each fungicide (pure and in the fungicide mixture) exhibited significantly different clearance rates. With some compounds, one or more stereoisomers inhibited the clearance of the other stereoisomers. Results were consistent across genders. In an effort to explain the observed stereoisomer clearance rates, we utilized a directed ligand-based pharmacophore analysis to identify key ligand features. Triadimefon was the only fungicide that preferentially underwent stereoselective carbonyl reduction rather than P450-mediated oxidation. Reduction of the prochiral carbonyl produced four stereoisomers of triadimenol. Relative formation of the triadimenol stereoisomers varied among 16 vertebrate species; the individual triadimenol stereoisomers differentially inhibited CYP3A4 metabolism. These results suggest that treating a chiral pesticide as a single chemical rather than a mixture could introduce errors in risk assessment. Although this work was reviewed by EPA and approved for publication, it may not necessarily reflect official Agency policy.

In the present in vitro study, it is demonstrated that the metabolite 4'-OH-deltamethrin is more cytotoxic than the parent compound deltamethrin. This work was supported by projects Refs. GR3510-A UCM-BSCH, S2009/AGR-1469 and Consolider-Ingenio CSD/2007/00063 (FUN-C-FOOD), Madrid, Spain.

### PS 593 Effect of Age on Plasma Protein Binding of Deltamethrin, cis-Permethrin, and Trans-Permethrin in Rats.

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Plasma protein binding (PPB) can influence the toxicokinetics of highly-bound chemicals, by limiting the amount of compound free to reach target organs and sites of elimination. Because the influence of age on PPB of common pyrethroids is unclear, this study was undertaken to determine if binding of deltamethrin (DLM), cis-permethrin (CIS) or trans-permethrin (TRANS) changes during maturation. The PPB of DLM, CIS and TRANS was studied in plasma of 10-, 15-, 21- and 90-day-old (adult) rats. A solvent extraction method was developed to quantify the binding of these highly hydrophobic compounds to rat plasma (RP) proteins and lipoproteins. A 10-μl aliquot of 14C-DLM, CIS or TRANS was mixed with 90 μl of plasma and cold compound yielding concentrations of 250-100,000 nM. Samples were incubated and shaken at 37°C for 3 hr. The unbound fraction was extracted with 200 μl of isooctane, the lipoprotein-bound fraction with 200 μl of octanol, and the albumin-bound fraction with 200 μl of acetonitrile. Samples were treated with 10 μl of a 0.64 M solution of sodium fluoride to inhibit carboxylesterases. The free fractions of DLM, CIS and TRANS were higher in 10-day-old RP (27.1 ± 6.1 to 29.9 ± 6.4 %) than adult RP (19.2 ± 1.5 to 21.3 ± 2.1 %). The fractions of DLM, CIS and TRANS bound to lipoproteins were slightly higher in the pre-weanlings, however the fractions bound to protein were substantially lower. In 15-day-old rats the free fraction and lipoprotein-bound fraction of DLM, CIS and TRANS decreased somewhat. At day 21, PPB reached adult levels. The lower PPB in pre-weanlings is consistent with lower total plasma protein levels.

The lower albumin levels appear to account for the relatively low protein binding of DLM, CIS and TRANS, (Supported by the Council for Advancement of Pyrethroid Human Risk Assessment).

### PS 594 Assessment of an In Vitro Human Dermal Absorption Database for Pesticide Formulations to Establish Scientifically-Based Default Dermal Absorption Values.

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Dermal absorption (DA) values for plant protection products (PPP) are required as inputs to risk assessment models used in the EU for assessing non-dietary risk for sprayers, workers, residents and bystanders. DA values are generated using undiluted PPP and representative field spray dilutions. The European Food Safety Authority (EFSA) published guidance on dermal absorption for PPP that includes highly conservative default DA values of 25% and 75% for undiluted PPP and field spray dilutions, respectively. Moreover, a default DA value of 75% applies to the undiluted PPP if it contains ≤5% active substance. To determine the validity of these default values, data from over 300 in vitro DA studies using human skin have been collected, collated and evaluated to establish the scientific validity of the default values and other endpoints. All studies that complied with the principles of OECD test guideline #428 were chosen for this analysis. These studies were performed on a wide range of formulation types and concentrations. The analysis was carried out considering two definitions of DA: (i) radioactivity present in receptor fluid and receptor chamber wash, plus skin minus the first two tape strips (i.e., worse-case definition) and (ii) radioactivity present in receptor fluid and receptor chamber wash plus skin minus stratum corneum (best-case definition). The analysis suggests that the default DA values for concentrate and spray dilution should not be more than 5% and 40%, respectively, considering the worse-case DA definition. The DA ranking for the different formulation types on DA-treated PPP is EC = FW > SL > SE > CC > CS = solid formulations > FS. Data will also be presented on the influence of formulation types on DA for field spray dilutions.
Pyrethroids are the more stable synthetic structural analogues of naturally occurring pyrethrins, which are potent insecticides that exhibit low mammalian toxicity. A reproducible and sensitive bioanalytical method was developed to monitor the uptake and elimination of pyrethroids in plasma and tissues following oral dosing of rats as part of an approach to assess their risk to human health by physiologically based pharmacokinetic (PBPK) models. Blood and tissue homogenates (100 μL) were spiked with Deltamethrin, cis-or trans-permethrin. 1% phosphoric acid (100 μL) was added (to reduce binding of pyrethroids to proteins) and mixed. Internal standard (another pyrethroids) and 400 μL acetonitrile were added and mixed (to precipitate the proteins and extract the pyrethroids) and centrifuged. Supernatant (500 μL) was evaporated to dryness, reconstituted with toluene (300 μL), filtered (0.2 μm PTFE syringe filter), evaporated, and reconstituted with toluene (50 μL) for analysis. Samples were injected into an Agilent 6890 gas chromatograph using pulsed splitless injection onto a Zebron® ZB5-MS capillary column eluting into a 5973 quadrupole mass analyzer in negative chemical ionization mode. Fragment ions were monitored using selected-ion monitoring for quantitation and verification of the analyte. The method was linear from 1 ng/mL to 1 μg/mL from tissues and blood. The limit of detection was 0.5 ng/mL, the limit of quantitation was 1 ng/mL from tissues

Parent insecticide residues dissipate on sprayed leaf surfaces and may yield derivations used as biomarkers of exposure to the pesticide on foliar surfaces. It is imperative to determine if surface derivatives actually contribute to the apparent aggregate exposure of harvesters based upon urinary biomonitoring. If pesticide derivatives that are potential urine biomarkers persist longer than the parent insecticide on treated plants and they are bioavailable, their absorption and excretion may confound harvester exposure reconstruction for risk assessment. Strawberry harvester exposures to malathion and fenpropatrin use were studied during July 2012 at Santa Maria, CA. Quantitative measurements of parent residues and corresponding biomarker residues on leaf surfaces determined using nitrile gloves worn by workers and 24 hr urine samples from harvesters (and matched controls with no occupational exposure) were conducted at specified intervals post malathion and fenpropatrin application on strawbery fields. Urine biomonitoring allows reconstruction of absorbed dose/day from all possible routes of pesticide and biomarker exposure using rapidly excreted urine biomarkers. Preliminary data show that when foliar malathion residue transferred to cotton cloths (μg/cm²; 0.238 to 0.027) and harvester gloves (μg/pair; 1394 to 3435) declined by 90% and 75% respectively in the 3d between the first picking and the second picking; comparing malathion transferred foliar biomarkers dropped only by 16% and 11%, respectively. These data suggest surface derivatives last longer than parent residues on foliar surfaces. Structurally similar derivatives are dermally absorbed and this may explain why biomonitoring historically has indicated exposure to parent long after the parent has dissipated.

**Toxicokinetics of Low Doses of the Pyrethroid, Deltamethrin (DLM), in Rats.**

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Pyrethroid insecticides are the widely used to control a wide variety of pests in and around homes, in food handling establishments, in mosquito control, and in agriculture. Pre-weaning and weaning rats are much more sensitive to lethality from a single high dose of the Type II pyrethroids deltamethrin (DLM) and cypermethrin than are adult rats. The hypothesis is that this is due to the incomplete development of the detoxification enzymes in the young rats. No age-related sensitivity to lethality or acoustic startle response has been noted at low doses of DLM. Our hypothesis is that, despite immature detoxification enzymes, sufficient metabolic capability is present to readily eliminate low doses. The current work was conducted to generate low-dose DLM blood and tissue time-course data to test this hypothesis and to provide input to a physiologically based pharmacokinetic model. Time-courses for 0.1, 1 and 5 mg DLM/kg have been delineated in adult rats dosed orally with DLM in 1 or 5 mL corn oil/kg or in 1 mL glycerol formal/kg. Serial sacrifices of groups of 3–5 rats have been conducted at intervals from 15 min to 96 hr post-dosing to obtain blood, brain, and other tissue samples for GC-MS analysis. Dose-dependent increases in peak DLM blood and non-adipose tissue levels were manifest from 2 to 6 hr. The larger volume of corn oil resulted in a relatively broad peak or plateau in blood levels and a more gradual decrease over the monitoring period than with the low volume. Brain levels at comparable times were significantly lower with the larger dosing volume. A significant portion of each dose of DLM was sequestered in fat, with slow release resulting in prolonged elevation of blood levels and thus availability to other tissues. (Supported by the Council for the Advancement of Pyrethroid Human Risk Assessment).

**Soil Concentrations and Human Exposures to Contaminated Soil in the Remote First Nation Community of Fort Albany, Ontario, Canada.**

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A study was conducted in the James Bay Region of Ontario, Canada to examine the concentration of dichlorodiphenyldichloroethane and metabolites (o,p'-DDE, p,p'-DDE, o,p'-DDD, p,p'-DDD, p,p'-DDT, p,p'-DDT, and ΣDDT) in potentially contaminated soil sites, and to assess the exposure pathways to ΣDDT by comparing the estimated daily intake (EDI) to acceptable daily intake (ADI) values. The contaminated soil plots were analyzed by a gas chromatograph equipped with a 63Ni electron capture detector (GC/ECD) or by gas chromatography with a mass spectrometer as a detector (GC/MS). The samples were first extracted using a Soxhlet apparatus with dichloromethane as the solvent and sample cleanup was applied using a Florisil packed column. The mean ΣDDT found in the soil plots were distributed irregularly with values ranging from not detected to 4.19 mg/kg. From the soil plots analyzed, Plot A had the highest ΣDDT concentration of 1.12 mg/kg, followed by Plot B and Plot C which were 0.09 mg/kg and 0.01 mg/kg, respectively. The exposure analysis showed that the risks to humans was below government guidelines, even though the DDT concentration in the soil was above Canada's soil guidelines of 0.7 mg/kg as set by the Canadian Council of Ministers of the Environment. The DDT concentration in Plot A breached maximum levels, therefore this land cannot not be used for agricultural as planned nor for recreational purposes. However, both Plots B and C were below maximum threshold limits, and this land may be used to grow safe food resources.

**Effect of Soil Fumigation with Methyl Bromide against Earthworm, Apopectrodea caliginosa.**


Methyl bromide, MBр, is a soil and food commodity fumigant that is used to control most kind of pests that born from or affect them, respectively. Earthworms, Apopectrodea caliginosa, are very important for healthy and productive soil. Although, MBр has been added to the list of chemicals that deplete the ozone layer (Montreal Protocol), its use still allowed under several condition worldwide. Effect of MBр against earthworm was investigated in this study. Worms were exposed to one fumigation level of 30 gm/m2 where worms were distributed in several depths
of the used soil. Mortality and some biochemical effects in the exposed worms com-
pared to controls were measured at 24, 48, and 72 hours after exposure. Higher MB at the
tested level caused 100% mortality in worms crawling at 10 and 30 cm depth in all tested intervals. At the 50 cm depth, mortality rates were 80, 90, and 90 percent at the 24, 48, and 72 hours after exposure, respectively. The effect of MB against protein content of earthworm crawling at the 50 cm depth was not recorded. Results showed that the MB increased significantly. There was no significant difference after 72 h of exposure in protein content between exposed worms and the control. Effect of MB at the tested level on the activities of earthworms glutathione-S-transferases, GS-T; acetylcholinesterase, AChE; and cellulase were also determined in worms crawling at the 50 cm depth. Results showed only slight significant increase in AChE activ-
ity at 24 and 48 hours after exposure relative to control. MB at the tested level caused significant increase in GS-T activity of the exposed earthworm at all tested exposure intervals. On the other hand, there was a significant increase in cellulase activity after 24 h as compared to control. However, this difference disappeared at 48 and 72 hours following exposure.

These results indicate that MB is a very potent soil fumigants that also kill the benign-
ciency earthworms in soil and protection of earthworms must be taken into con-
sideration while searching for alternative to MB.

**600 Mechanism of Paraquat-Induced Pulmonary Fibrosis and Intervention of Pyrrolidine Dithiocarbamate.**

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The mechanism of Paraquat-induced pulmonary fibrosis and potential therapeutic effect of pyrrolidine dithiocarbamate (PDTC) were studied. Male SD rats were di-
vided into control group (0.9% NaCl, gavage), PDTC group (100mg/kg, ip), PQ (80mg/kg, gavage) group and PQ+PDTC (100mg/kg, ip) group. On the1st, 3rd, 7th, 14th, 28th and 56th day after treatment, The expressions of connective tissue growth factor (CTGF) and α smooth muscle actin (α-SMA) in lung tissues were measured. The mRNA levels of CTGF, Fn, Col I and Col III and integrin α5 were an-
alyzed with quantitative RT-PCR. Meanwhile, the lung pathological changes were observed and the content of Hydroxyproline (Hyp) was measured. The expression of CTGF in PQ group increased gradually compared with control group (P<0.05). CTGF mRNA level significantly increased from the 1 st to the 14 th day compared with control group (P<0.05). PQ significantly increased Fn mRNA level on all time points and integrin α5 mRNA level from the 3 rd to 56 th day compared with control group (P<0.05). Col I mRNA level significantly increased from the 7 th to the 56 th day and Col III mRNA level appears to be decreased from the 14 th to the 56 th day. PDTC treatment significantly decreased the levels of those factors compared with PQ group in corresponding time points (P<0.05). Noteworthy, PDTC strongly attenuated histopathological changes and decreased the content of Hyp. These results suggested that CTGF plays a key role in paraquate-induced pul-
monary fibrosis, which is characterized by increased Fn, integrin α5, Col I and Col III mRNA levels. PDTC may inhibit NF-kB activity and further significantly de-
crease expressions of CTGF leading to drastically attenuated pulmonary fibrosis. However, the mechanisms of PDTC intervention still remain to be explored.

**601 Testicular Toxicity of Fluorochloridone in Adult Sprague-
Dawley Rats.**

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Fluorochloridone (FC) a widely used herbicide in Europe and the USA, has been rec-
ognized as a potential reproductive and developmental toxicant. However, there is little data available concerning its male reproductive toxicity. In this study, we ex-
amined the testicular toxicity of FC in SDW rats. Adult rats were treated 5x gav-
age with FC at doses of 0, 30, 150, 750 mg/kg/d for four weeks. FC exposure re-
sulted in a decrease in the absolute and relative weight of testes and a decrease in the absolute weight of epididymides as compared with the control. Cauda epididymal sperm count decreased dramatically in a dose-dependent manner. In addition, his-
tological lesions were also found in the testes of the treated animals. A dose-effect (response) relationship analysis suggested that changes in cauda epididymal sperm count and testicular histological structure could be the most sensitive indicators for FC induced testicular toxicity.

**602 Carbofuran Is Not an Endocrine Disruptor—Weight-of-
Evidence of US EPA’s Tier 1 Endocrine Disruptor Screening Program Assays and Higher Tier Studies.**

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Carbofuran, the active ingredient in FMC Corporation’s FURADAN® Insecticide/ Nematicide products, underwent testing in EPA’s Tier 1 endocrine dis-
ruption screening program (EDSP), as required for all US registered pesticide active
ingredients, to assess whether carbofuran had the potential to interact with en-
docrine systems. Nine of the eleven carbofuran Tier 1 EDSP assays showed no po-
tential to interact with endocrine systems. The slight increase in estradiol in the in
vitro steroidogenesis assay, and the mild to moderate thyroid hyperplasia and mild
hyperplasia in the amphibian metamorphosis assay observed with relatively high
concentrations were not corroborated by any of the other assays. Only the highest concentration (100 µM) was positive in the steroidogenesis assay, several orders higher than the concentration required for AChE inhibition. Previously conducted higher tier in vivo carbofuran developmental, reproductive and chronic toxicity studies showed no endocrine-mediated effects. The overall lack of effects in the carbofuran Tier 1 EDSP assays is consistent with the lack of en-
docrine-mediated effects in the previously conducted carbofuran higher tier in vivo studies. The collective results of the carbofuran Tier 1 EDSP assays and previously conducted higher tier in vivo studies indicate carbofuran is not an endocrine dis-
ruptor, and that there is no concern for endocrine effects resulting from the use of FURADAN® products according to label instructions.

**603 Reproductive Outcomes in Women Para-Occupationally Exposed to Pesticides in an Agricultural Community in Southern Mexico.**

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Epidemiological studies have shown that farmers are at high risk of developing ad-
verse effects including reproductive problems. However, reproductive outcomes by para-occupational exposure to pesticides in waves of farmers are poorly docu-
mented. We reported reproductive alterations such as poor semen quality in farm-
ers and spontaneous abortions and preterm birth in waves of agricultural workers
mostly exposed to organophosphate pesticides (OP). Thus we evaluated the scenery of para-occupational pesticide exposure and its associated reproductive outcomes in waves of farmers. A transversal study was conducted in Yucatan, Mexico in waves of farmers exposed to pesticides (n=28) and waves of workers without pesticide expos-
ure (n=45). Women donated a blood sample and responded a questionnaire. The TORCH profile was determined as potential confounders. General characteristic and TORCH profile were similar between two groups. Age at menarche was margi-
ally lower in waves of workers (11.5 ± 1.06 vs. 12.13 ± 1.56 years, p=0.07).

Difficulty getting pregnant was more prevalent in wives of farmers (63 vs 13%, p<0.0003), preterm births were marginally more frequent in these women (7 vs 0%, p=0.07), and the birth weight of the first child was slightly lower in couples of farmers (2.94 ± 0.55 vs 3.16 ± 0.48 Kg, p<0.09). A complex scenery of pesticides ex-
posure was observed, parathion, glyphosate, 2,4-D amine, methamidophos and chloropyrifos were among the most used pesticides. Results show that pesticide para-
occupational exposure in waves of farmers may be involved with reproductive out-
comes. Supported by PROMEP-SEP-México.

**604 Extension of a Nasal Dosimetry Model for Acetaldehyde to Account for Vasodilation.**

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Acetaldehyde (AAl) is an important industrial chemical, used in manufacturing a wide range of products, but sources also include volcanoes, forest fires, biological waste degradation, and respiration by plants. Subchronic inhalation causes olfac-
itory and nasal respiratory tissue degeneration in rats at ≥ 150 ppm and nasal tu-
mors in both regions at ≥ 750 ppm. The toxicity of AAl is likely due to its metab-
olism to acetic acid in nasal tissues. A series of physiologically-based
pharmacokinetic (PBPK) models have been published which result from computational and experimental data to specify aerial and air/phase transport resistance in the nose. Most recently, Teegarden et al. (Inhal. Toxicol., 28:375-390, 2008) developed a version which included polymorphisms in human high-affinity acetaldehyde dehydrogenase (ALDH2). However, Stanek et al. (Inhal. Toxicol., 13:807-822, 2001) showed that ALDH causes vasodilation, increasing uptake of acetone with co-exposure of the two. Vasodilation will also increase uptake of AlAD. If this is not included in the model, the extent of metabolism estimated by fitting a PBPK model to nasal AlAD uptake data could be overpredicted at concentrations where dilution is significant. Therefore we revised the PBPK model of Teegarden et al. to include vasodilation (increases in nasal blood flow), with the amount of dilution estimated from the data of Stanek et al. using a parallel acetone PBPK model. An error in model code calculation of tissue phase mass transfer resistance was also corrected. The net result is an increase in the estimated Vmax for ALDH2 and a dose-dependent change in total metabolism in rats: increased at lower concentrations where ALDH2 is not saturated, but decreased at 1500 ppm where vasodilation is significant. Quantifying the impact of this revision requires integration of model results with dose-response (i.e., benchmark dose) modeling. (Views expressed here are those of the authors and do not necessarily reflect the views or policies of the US EPA.)

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**605 An In Silico Model of Spermatogenesis for Use in Predictive Toxicology.**

EPA’s toxicity reference database (ToxRefDB), covering 963 compounds tested in 4209 animal studies, indicated testicular atrophy as an adverse outcome for 278 (28.9%) of these chemicals. Several compounds have been tested in chronic and subchronic rodent toxicity studies. The sensitivity of the male reproductive system, coupled with the biological complexity of testicular function and the large number of untested chemicals, motivates the need for predictive computational models of male reproductive function based on high-throughput screening (HTS) and in vitro data. Toward this end, we are building a virtual tissue model of the hyperemic-putative-germinal system (hHPGS) that integrates computational approaches for kinetic modeling and dynamic simulation. As a proof of concept, we have constructed a multicellular agent-based model of the rat seminiferous tubule using CompuCell3D. The model integrates cell signaling networks for hormonal effects (FSH, testosterone), growth factor and cytokine stimulation (GDNF, TGFβ, IL1β) and transcription factors (e.g. ERM, Rho5, SOX8, WT1) responsible for regulating key events in the spermatogenic cycle. It also enables Sertoli cell interactions including cytoskeletal restructuring and protein secretion. To test our model, over 1900 chemicals associated with toxic effects on sperm were characterized for effects on spermatogenesis and mechanisms of action. We then identified chemicals targeting specific key aspects of spermatogenic cell division (bleomycin, cisplatin, doxorubicin), cell/cell adhesion (adjudin, indenopyrindinates, indazole-3-carboxylic acid), and Sertoli cell death (β-benzene hexachloride), as well as putative endocrine disruptors (e.g., atrazine, bisphenol A, ketocortazole). Preliminary results indicate that the model responds appropriately, in terms of sperm production, to the disruption of key signaling events. This abstract does not necessarily reflect Agency policy.

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**606 Development of PBPK Models for Gasoline in Adult and Pregnant Rats and Their Fetuses.**

Concern for potential developmental effects of exposure to gasoline-ethanol blends has grown along with their increased use in the US fuel supply. Physiologically-based pharmacokinetic (PBPK) models for these complex mixtures were developed to address dosimetric issues related to selection of exposure concentrations for in vivo toxicity studies. Sub-models for individual hydrocarbon (HC) constituents were first developed and validated with published PBPK data where available. Successfully calibrated sub-models for individual HCs were combined, assuming competitive metabolic inhibition in the liver, and a priori simulations of mixture interactions were performed. Blood HC concentration data were collected from exposed adult non-pregnant (NP) rats (9K ppm total HC vapor, 6h/day) to evaluate performance of the NP mixture model. This model was then converted to a pregnant (PG) rat mixture model using gestational growth equations that enabled a priori estimation of life-stage specific kinetic differences. To address the impact of changing relevant physiological parameters from NP to PG, the PG mixture model was first calibrated using the NP data. The PG mixture model was then evaluated using data that were subsequently exposed (9K ppm/6.33h gestation days (GD) 9-20). Overall, the mixture models adequately simulated concentrations of HCs in blood from single (NP) or repeated (PG) exposures (within -2-3 fold of measured values of most HCs), indicating that the blood data from PG rats were not highly sensitive to PG-specific physiological parameters. This PG model mimics external HC concentrations in the total fetus during in utero exposure to HC vapors. This is an abstract of a proposed presentation and does not necessarily reflect EPA policy.

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**607 Strategy to Support Clearance Model of Inhaled Libby Amphibole Asbestos Fibers in Rat and Human Respiratory Tract.**
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To characterize inhaled Libby amphibole (MA) asbestos fibers as retained dose (RD), we developed a strategy to integrate experimental data with compartmental modeling. Compartment extend our model structure for inhalability and deposition (Asgharian, 2012) to estimate RD in major respiratory tract (RT) regions (upper respiratory tract, URT; tracheobronchial, TB; and pulmonary, PU), pleural lining (PL) and lymph nodes (LN). The strategy provides data to derive 3 clearance models of the respiratory tract (RT): mucociliary (MC), transcytosis (TR), and dissolution (DS). Initial mass in each region is calculated using the deposition model verified with fiber burden data in the URT, trachea/larynx, 5 lung lobes and pleural casts in F344 rats from a 6-hr exposure of LA at 0, 3.5, or 25 mg/m³. MC rates of the URT and TB regions are estimated by refining published rate constants with fitting mass burdens measured in URT, TB, LN and PL compartments of a 5-d study in rats at those 3 concentrations with post-exposure time course (0-, 6-, 12-, and 24-hr). Burden data in rats from a 13-wk study at 0, 3.3, 1, and 10 mg/m³ with post-exposure time course (1d and 1-, 3- and 18-mo) refine TR rates from PU to TB, PL and LN compartments and evaluate overload. The initial DS rate is based on burden data from the URT considered in vitro data on DS of LA incubated with synthetic lung lining fluid. Data from in vitro incubation of LA with acid refine DS rate for TB and PU regions. This integrated strategy is the first to derive characterization of MC, TR and DS rates in the rat RT. For humans, DS rates were not scaled, published human MC rates are adjusted based on mass conserva- tion (Asgharian et al., 2001). TR rates to PL and LN compartments are scaled on MC, and PU TR is scaled from the rat by an approach described for particles. (Jarabek et al., 2005) based on regional surface area. (The views expressed in this abstract are those of the authors and do not necessarily represent the views or policies of the U.S. Environmental Protection Agency.)

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**608 A Modified PBPK Model for RDX (hexahydro-1, 3, 5-trinitro-1, 3, 5-triazine) to Improve Rat to Human Toxicokinetic Extrapolation.**

RDX is a military explosive that has been detected in air, soil, and ground water at or near military bases and munitions plants and storage facilities. Studies in rats provide evidence of mortality, reproductive toxicity, and neurotoxicity associated with RDX exposure. Physiologically-based pharmacokinetic (PBPK) models can aid in interpreting toxicological data and extrapolations across dose, species and exposure routes. Presented here is the further development and application of an RDX PBPK model for rats and humans originally published by Sweeney et al. (Regul. Toxicol. Pharmacol., 2012 x: 72, p. 107-114). The differences in the GI tract submodel and absorption parameters for different formulations of RDX in the Sweeney model are problematic for risk assessment because there is no rationale or predictability to the changes. Modifications to the model include a consistent submodel for the GI tract, a consistent set of absorption parameters to simulate oral exposure to RDX, and oral absorption rate constants keyed to different formulations of administered RDX per the kinetic data in the literature. Model code was also added to simulate drinking water and inhalation exposures relevant to human chronic exposures. The resulting model simulations are consistent with observed differences in the time course blood level data from a variety of PK studies, RDX formulations, and dosing regimens. Blood concentrations of RDX did not differ considerably with simulating inhalation exposure as compared to oral exposure. Model fits to rat data resulting from a subchronic drinking water exposure demonstrated achievement of steady state levels, lack of metabolic induction over time.
Developing New Methods to Measure Reactive Oxygen Species Related to Aerosol Components.

W. Huyyn1, E. Brown2 and R. E. Pelizzari, 1Public Health - Environmental Health Sciences, University of Massachusetts Amherst, Amherst, MA; 2Veterinary & Animal Sciences, University of Massachusetts Amherst, Amherst, MA.

Reactive oxygen species (ROS) have been shown to be an important indicator of adverse toxicological effects and has been widely studied in the context of ambient air pollution exposure research. ROS is formed in vivo as a result of exposure to ambient air pollutants and is thought to be a convenient approach to understanding responses arising from air pollution exposures. Current methods of measurement for ROS typically follow a labor-intensive and time-consuming method of collection and extraction of particles from the field, which are then introduced (often at very high concentrations) in vitro. The objectives of this research are to streamline these approaches by developing and characterizing a new instrument capable of automated, semi continuous quantification of ROS. The work presented here attempts to fill this gap by repeatedly inducing reactive oxygen on plated immortalized pulmonary epithelial cells (BEAS-2B) which have been loaded with 2',7'-dichlorofluorescein diacetate (DCFH-DA). The data presented here are based on a method using 96-well plates and standard fluorescent spectrometry methods. These results are compared with similar approaches using a system of horseradish peroxidase and DCFH-DA. Tested elements include micromolar concentrations of hydrogen peroxide, solutions of ammonium sulfate and nitric acid, and quinones. We show that, with notable limitations, plated cells can respond to repeated dosing of simulated air pollution, suggesting that this method is suitable for further instrument method development.

Designing Quantitative Structure Activity Relationships (QSAR) to Predict Specific Toxic Endpoints for Polybrominated Diphenyl Ethers (PBDE) in Mammalian Cell Culture Systems.

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Polybrominated diphenyl ethers (PBDEs) are flame retardants that have had vast industrial application in consumer products, such as plastics, building materials, electronics and textiles. They are structurally similar to thyroid hormones that are responsible for regulating metabolism in the body. Therefore, PBDEs compete for the thyroid hormone binding receptors and this can adversely affect thyroid hormone transport and metabolism. Due to their potential threat to human health, this study aimed to design Quantitative Structure Activity Relationships (QSAR) models for predicting specific toxic endpoints, namely, cell viability and apoptosis. Human hepatocarcinoma (Hep G2) cells were exposed to PBDEs and were used as a model system to evaluate cell viability using Janus Green dye and apoptosis using a caspase assay. Data collected from the experiments were used to create QSAR models using the Genetic Function Approximation (GFA) method of generating models for predicting specific toxic endpoints, namely, cell viability and apoptosis. Human hepatocarcinoma (Hep G2) cells were exposed to PBDEs and were used as a model system to evaluate cell viability using Janus Green dye and apoptosis using a caspase assay. Data collected from the experiments were used to create QSAR models using the Genetic Function Approximation (GFA) method of generating predictive models. Cell viability and apoptosis responses elicited by the PBDEs models using the Genetic Function Approximation (GFA) method of generating models for predicting specific toxic endpoints, namely, cell viability and apoptosis. This study aimed to design Quantitative Structure Activity Relationship (QSAR) models for predicting specific toxic endpoints, namely, cell viability and apoptosis. These results are compared with similar approaches using a system of horseradish peroxidase and DCFH-DA. Tested elements include micromolar concentrations of hydrogen peroxide, solutions of ammonium sulfate and nitric acid, and quinones. We show that, with notable limitations, plated cells can respond to repeated dosing of simulated air pollution, suggesting that this method is suitable for further instrument method development.

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hour (AML) - for two exposure scenarios (single 0.25 liter drink of water or 10 minute shower) under typical (10 ppb) and plausible high level (50 ppb) water concent-
trations. MMPGGL and LW (as a fraction of body weight) values used in the analysis reflect the range of values reported for adult humans. For each concent-
tration, each dose metric was changed less than 5% for the showering scenario (in-
halation and dermal exposure) because Vmax for hepatic metabolism is not a very influential parameter for dose metrics related to blood concentration following low
level exposures. In contrast, an 8-fold difference in Cmax and AUC was observed for each oral exposure concentration, but AML was relatively unchanged. Sensitivity analysis for AUC for the oral exposure scenario revealed that MMPGGL was the most influential parameter, followed closely by LW, with blood flow to the liver being moderately influential. This analysis demonstrates that variability in the scaling factors used for in vitro to in vivo extrapolation (IVIVE) of metabolic rate parameters can have a significant impact on estimates of internal dose metrics under environmentally relevant exposure scenarios. This indicates the need to evalu-
ate both uncertainty and variability for scaling factors used for IVIVE. (This abstrac-
t does not necessarily reflect USEPA policy).

614 Kinetic Modeling Reveals the Roles of ROS Scavenging and DNA-Repair Processes in Shaping the Dose-Response Curve of KBrO3-Induced DNA Damage.


We have used kinetic modeling to investigate how DNA repair processes and scav-
engers of reactive oxygen species (ROS) can affect the dose-response shape of pro-
oxidant induced DNA damage. We used an example chemical KBrO3, a water
ozonation by-product and environmentally present pro-oxidant with genotoxic and
carcinogenic effects. In our model, KBrO3 is activated via interaction with glu-
thathione and forms reactive intermediates that directly interact with DNA to form
8-hydroxy-2-deoxyguanosine DNA adducts (8-OH-dG) – an effect convincingly
established in literature. The single strand breaks (SSB) that can result from failed
base-excision repair of these adducts were considered in the model as an effect
downstream from 8-OH-dG. We previously demonstrated that in the presence of
effective base-excision repair, 8-OH-dG can exhibit threshold-like dose-response
dependence, while the downstream SSB can exhibit a linear dose-response. We
now further demonstrate that this result holds for a variety of model variants. In
particular, we investigated how the presence of a scavenger of the bromate reactive
intermediates affects the dose-response shape of 8-OH-dG and SSB. It has been
shown that melatonin, a terminal antioxidant, inhibits KBrO3-caused oxidative
damage. Our modeling revealed that a single pulse exposure to KBrO3 in the pres-
ence of such a terminal scavenger can lead to a sublinear/threshold-like dependence
of the response for both 8-OH-dG and SSB. However, sustained exposure to
KBrO3 can lead to fast scavenger exhaustion, in which case the dose-response
shapes for both endpoints are not substantially affected. The results are important
to consider when forming conclusions on a chemical's toxicity based on the dose-
response of early genotoxic events.

615 A Compartmental Pharmacokinetic (PK) Model for Bromate in F344 Rats.

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Bromate (BrO3-) is a toxic water disinfection by-product formed during ozonation of source water containing bromide (Br-). It is a proven animal and probable human carcinogen (group 2B). However, BrO3- pharmacokinetics (ADME) at low
doses is not fully understood. To better understand BrO3- ADME, we developed a pharmacokinetic (PK) model based on the raw data from our recent publication (Bull et al. 2012). Based on goodness of fit, and the model diagnostics, a pharma-
cokinetic model fitted with individual animal data and weight-2 (1/WT2) weight scheme was chosen as a model of choice to describe BrO3- disposition in female F344 rats. BrO3- disposition after IV bolus was best described using a 1-compartmental pharmacokinetic model for doses up to 0.5 mg/kg; and by using a 2-compartmental pharmacokinetic model for doses of 1-2.5 mg/kg potassium bromate (KBrO3). BrO3- disposition was best described using a 1-compartmental model follow-
ing its oral administration (0.5 to 20 mg/kg KBrO3) in female F344 rats. Analysis of BrO3- PK parameters following oral administration, suggested that BrO3- absorption occurs in a first order manner with a rate constant (Ka) of ~0.16
min-1. BrO3- appears to undergo extensive first pass reduction, resulting in
bioavailability of ~19-25%. Approximately 90% of orally administered BrO3- is re-
duced to Br- in the gut, and BrO3- and its remaining BrO3- appear to be excreted from the body as BrO3- in the urine at 0.002 L/min/kg. In conclusion, BrO3- undergoes
extensive reduction to Br- in liver and blood following its oral administration in
female F344 rats, which limits BrO3- distribution to the peripheral tissues to exert its
toxic effects.

616 A Strategy for Developing a PBPK Model to Describe the Kinetics of Silver Nanoparticles.

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Our previous model for C60 was proposed as a general platform for PBPK model-
ing of engineered nanoparticles (NPs). Since the structure was based on biological
mechanisms that govern NP disposition, the platform can be applied to other NPs
by changing parameters to account for particle-to-particle variance in biological
processes. In this study, the model was parameterized for silver (Ag) NPs using tis-
sue concentration data collected in adult female SD rats after a single intravenous
(IV) dose (1 mg/kg) of Ag NPs (20 or 110 nm) in water stabilized with polyvinyl
pyrrolidone (PVP) or Ag acetate in PVP. Parameterization was based on size-de-
dependent dissolution potential of Ag NPs under physiological conditions as well as
generally applicable size-dependent endocytosis efficiency for particles. Due to the
dissolution potential, a combination of two kinetic models, one for Ag NPs and the
other for ionic forms of Ag was used. The model successfully simulated observed
concentrations of total Ag in various tissues as well as in urine and feces after a
single IV dose of Ag NPs. The model was then extrapolated to Ag NPs with other
coating agents. The published data for tissue concentrations of total Ag in adult
male Wistar rats after single or multiple (~ 0.1 mg/kg/day) IV administration of Ag
NPs in phosphate buffer (20 or 110 nm) was simulated. For this evaluation, the im-
 pact of different coatings on Ag NPs stability in size, shape and surface properties
was also considered in the parameterization. The current model captured the time
courses of total Ag concentration in blood, liver, lung, spleen, and heart both for
single and multiple exposures from this study. The overall success in extending the
C60 model to Ag NPs demonstrated the applicability of our model structure, as a
general platform, to other NPs when combined with nanoparticle-specific proper-
ties for the given material (This work was supported by NIEHS Award #U19ES019525, but solely expresses the view of the authors).

617 A Physiologically-Based Pharmacokinetic Model to Describe the Pharmacokinetic Disposition for Hexamethyldisiloxane, a Linear Volatile Methyl Siloxane, following Inhalation Exposures in the Rat.


The purpose of this work was to expand a previously developed physiologically
based pharmacokinetic (PBPK) model for hexamethyldisiloxane (HMDS) (Dobrev et al., 2003) with additional kinetic data to further examine the processes regulating the pharmacokinetic disposition of HMDS. Time-course concentrations of parent HMDS and total metabolites in multiple tissues, blood and exhaled
breath in male and female F344 rats during and following nose only vapor inhala-
tion exposure to 5000ppm 14C-HMDS were used to extend the model. A 15 day
exhalation study was also conducted to understand impacts, if any, following re-
peated exposures. Based on similarities in physicochemical properties, the HMDS
model structure was adapted from another volatile methylsiloxane (VMS) oc-
tamethylcyclotetrasiloxane (D4) model structure (Andersen et al., 2001). This
updated model now includes saturable metabolism in the liver, a diffusion-limited
uptake of HMDS in the fat compartment and a blood lip pool where a fraction of
HMDS is sequestered and unavailable for exchange into blood and exhalation from
the lung. HMDS and its metabolites in blood, liver, lung, fat and exhaled breath
following both single and multiple exposures are well described with this model
using a single set of parameters. Consistent with D4 kinetic behavior, low blood/air
partitioning and extensive metabolism lead to rapid clearance of free HMDS from
the body following both single and repeated exposures. Despite retention of bound
HMDS in tissues at the extended time points following exposure, the extensive and
rapid clearance mechanisms of HMDS lead to tissue and exhaled breath concentra-
tions of HMDS that are similar following both multiple day exposures and single
exposures. This PBPK model is a starting point from which the pharmacokinetic
disposition of HMDS and, upon extension, other linear VMSs, can be evaluated for
risk assessment.
Mathematical modeling has become an important tool to assess xenobiotic exposure in humans. In the present study, we have used a human physiologically-based pharmacokinetic (PBPK) model to reproduce the time-course of 3-hydroxybenzo(a)pyrene (3-OHBaP) in the urine of industrially exposed workers and in turn predict most plausible exposure scenarios. Urinary voids from a dozen workers highly exposed to polycyclic aromatic hydrocarbons (PAHs) in the Rhone-Alpes region in France have been collected during a typical workweek (beginning and end-of-shift) and subsequent days off; urinary concentrations of 3-OHBaP were then determined. Based on the information obtained for each worker (airborne BaP concentration, daily shift hours, tasks, protective equipment), the time courses of 3-OHBaP in the urine of the different workers have been simulated using the PBPK model, considering the various possible exposure routes, oral, dermal and inhalation, as well as combined exposure. The model was constructed from in vivo experimental data in rats and then extrapolated from animals to humans after assessing and adjusting most sensitive model parameters as well as species specific physiological parameters. The model was able to closely reproduce the observed time course of 3-OHBaP and establish most plausible exposure route depending on the worker. It appears as a useful tool to better interpret biomonitoring data of PAH exposure on the basis of 3-OHBaP biomarker levels.
levels. MCMC analysis was employed to infer population and individual distributions for these significant parameters based on levels measured in 12 individuals exposed to styrene, n-hexane or toluene. The generic VOA PBPK model calibrated by MCMC analysis was able to predict end-exhaled air levels for the 12 individuals and 3 compounds. The chains from the converged MCMC runs will be used to derive distributions of end-exhaled air for given exposure scenarios. Our next steps will be to include more individuals, compounds and exposures scenarios with variable workloads to better describe variability in the population.

623 Development of a New 3D Human Small Intestinal Tissue Model.
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The epithelial lining of the gastrointestinal (GI) tract is a gatekeeper for entry of orally ingested nutrients and xenobiotics including medications. The intestinal lining has a well organized structure containing proliferative cells which migrate along the crypt-villus axis and differentiate into functional mature epithelial cells. Currently, the most common in vitro model utilized for study of drug absorption at the intestinal mucosa is based on a 2-D cell culture model of the colon carcinoma cell line, Caco-2. These cells differentiate into monolayers of polarized enterocytes that are connected by tight junctions, but lack mucus-secreting goblet cells and show inter-passage inconsistency. Others have developed small intestinal organoids which are not suitable for apical application of test articles. Here, we report the reconstruction of a human organotypic small intestinal tissue (SI) generated from primary human SI epithelial cells that grow in tissue culture inserts using serum free medium. Human SI epithelial cells and myofibroblasts were expanded in monolayer culture and seeded on a microporous membrane containing a fibroblast-collagen-gel substrate to reconstruct three dimensional organotypic SI tissues. Tissue morphology and biomarker expression of the SI model were characterized by H&E staining and immunohistochemistry, respectively. Basal cell proliferation markers, cytokeratins (CK) and mucus were monitored over an 11-day culture period. Analysis of the SI tissue model revealed: 1) wall-to-wall growth of the epithelial layer, 2) columnar epithelial cell morphology similar to that of native SI tissue, 3) expression of muc-2, CK19, and villin at the surface of the epithelium, and 4) Lgr5+ (crypt stem cell marker) positive cells. In conclusion, the new human cell based 3D SI tissue model will likely serve as a valuable tool to evaluate pre-clinical therapeutic drug candidates intended for oral administration and study microbiota and microbial infection of the GI tract.

624 Evaluating Dose-Response Parameters of an In Vitro DNA-Chemical Interaction Using a Mechanistic Biological Model.
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Benchmark dose modeling of genomic data is a trending approach in toxicity testing, and system biology modeling attempts to understand quantitative data in light of what we expect from more complex biological systems. Biological modeling approaches can incorporate different forms of kinetic and dynamic data of a chemical interaction in vitro. The objective of this effort was to interpret a biological model mechanism and parameters describing a chemical doubled stranded oligonucleotide interaction assay for a series of chemicals (tetrachlorohydroquinone, styrene-7,8-oxide, methyl methanethiolate, phenyl glycidyl ether, and benzo(a)pyrene diol epoxide), with both time series and dose-ranged measurements. The chemicals have been analyzed with time series, dose-response and inhibition mixture response analysis. A model using mixed Michaelis-Menten kinetic and multiple binding was determined to be best suited with the data coherency between the biological and mathematical expectations. This was based on the results of the model, as well as the poorer results of other considered models (e.g. constant rate constants, linear rate constants, multi-step reactions, Hill functions). Correlations between coefficients and various chemicals, as well as sensitivity analysis, gave evidence of a deeper mechanism underlying nucleotide perturbations (e.g. relations amongst the half-life, dose and the curvature of the reactions). Two model parameters, molecular binding coefficient (range from 0.1 to 2.6) and nonlinear affinity rate constant (ranging from 30 to 5300 μM), were dose dependent and specific to each chemical. These kinetic parameters are suitable candidates for a quantitative structure-activity analysis with chemical descriptors for nucleotide binding sites and interaction efficacy as indicators of adfuct formation. These biological models fit the biological expectations for the governing mechanisms, giving further evidence of the effectiveness of statistical and benchmark dose modeling in toxicity testing.

625 Developing Predictive Approaches to Characterize Adaptive Responses of the Reproductive Endocrine Axis to Aromatase Inhibition: Computational Modeling.

Exposure to endocrine disrupting chemicals can affect reproduction and development in both humans and wildlife. We developed a mechanistic mathematical model of the hypothalamic-pituitary-gonadal (HPG) axis in female fathead minnows to predict dose-response and time-course (DRTC) behaviors for endocrine effects of a well-defined aromatase inhibitor, fadrozole (FAD). The model includes a regulatory feedback that mediates adaptive responses to endocrine stress by controlling the secretion of a generic gonadotropin (LH/FSH) from the hypothalamic-pituitary complex. Plasma 17β-estradiol (E2) and ovarian cytochrome P450 (CYP) 19A aromatase mRNA data from two time-course experiments, each of which included both an exposure and a depuration phase, and plasma E2 data from 4 days exposure experiment were used to develop and evaluate the model. Model parameters were estimated using E2 concentrations for 0, 0.5, and 3 μg/L FAD concentrations, and good fits to these data were obtained. The model accurately predicted CYP19A mRNA fold changes for controls and three FAD doses (0, 0.5, or 3 μg/L), and venous E2 dose-response during FAD exposure on day 4. Comparing the model-predicted DRTC with experimental data provided insight into how the feedback control mechanisms embedded in the HPG axis mediate these changes: adaptive changes in plasma E2 levels occurring during exposure and “overshoot” occurring post-exposure. This study demonstrates the value of mechanistic computational modeling to examine and predict the possible dynamic behaviors. This abstract does not necessarily reflect US Environmental Protection Agency policy.

626 Mitochondrial Dysfunction-Induced by Sertraline, an Antidepressant Agent.
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Sertraline, a selective serotonin reuptake inhibitor (SSRI), has been used for the treatment of depression. Although it is generally considered safe, cases of sertraline-associated liver injury have been documented; however, the possible mechanism of sertraline-associated hepatotoxicity is entirely unknown. Here we report that mitochondrial impairment may play an important role in liver injury induced by sertraline. In mitochondria isolated from rats, sertraline uncoupled mitochondrial oxidative phosphorylation and inhibited the activities of oxidative phosphorylation Complexes I and V. Additionally, sertraline induced Ca2+-mediated mitochondrial permeability transition (MPT), and the induction was prevented by bongkrekic acid, a specific MPT inhibitor targeting adenine nucleotide translocator (ANT), implying that the MPT induction is mediated by ANT. In freshly isolated rat primary hepatocytes, sertraline rapidly depleted cellular ATP and subsequently induced lactate dehydrogenase (LDH) leakage; both were attenuated by bongkrekic acid. Our results, including ATP depletion, induction of MPT, inhibition of mitochondrial respiration complexes, and uncoupling oxidative phosphorylation, indicate that sertraline-associated liver toxicity is possibly via mitochondrial dysfunction.

627 Mouse Liver Protein Sulphydryl Depletion Induced by Acetaminophen Exposure.
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Since Acetaminophen (APAP) remains the leading cause of acute liver failure in the western world, a large body of research has been conducted to understand the mechanisms behind the pathogenesis. The role of protein sulphydryl depletion in APAP-induced liver injury was investigated in this study and compared to protein adducts and classical measures of toxicity. A single oral gavage dose of 150 or 300 mg/kg APAP in B6C3F1 mice produced increased serum alanine aminotransferase, bilirubin, liver necrosis and glutathione depletion in a dose-dependent manner. The levels of global free protein sulphydryls, were significantly decreased at 1 hour and...
Acetaminophen (APAP) overdose is the most frequent cause of acute liver failure in the western world. Controversy exists regarding the hypothesis that the hepatocyte injury is amplified by a sterile inflammatory response, rather than being the result of intracellular mechanisms alone. A recent study suggested that the purinergic receptor antagonist A438079 protects against APAP-induced liver injury by preventing the activation of the Nalp3 inflammasome in Kupffer cells and thereby preventing inflammatory injury. To test the hypothesis that A438079 actually affects the intracellular signaling in hepatocytes, C57BL/6 mice were treated with APAP (300 mg/kg) and A438079 (80 mg/kg) or saline and GSH depletion, protein adduct formation, c-Jun-N-terminal kinase (JNK) activation, oxidant stress and liver cell necrosis was determined between 0-6h after APAP administration. APAP caused rapid GSH depletion, extensive protein adduct formation in liver homogenates and in mitochondria, JNK phosphorylation and mitochondrial translocation of phospho-JNK within 2h, oxidant stress, and extensive centrifugal necrosis at 6h. A438079 significantly attenuated GSH depletion, which resulted in a 50% reduction of total liver and mitochondrial protein adducts and substantial reduction of JNK activation, mitochondrial P-JNK translocation, oxidant stress and liver injury. The same results were obtained using primary mouse hepatocytes. A438079 did not directly affect JNK activation induced by tert-butyl hydroperoxide and GSH depletion. However, A438079 dose-dependently inhibited hepatic P450 enzyme activity. Thus, the protective effect of A438079 against APAP hepatotoxicity in vivo can be explained by its effect on metabolic activation and cell death pathways in hepatocytes without involvement of the Nalp3 inflammasome.

The hepatotoxicity noted was also apparent from the clinical chemistry results that showed marked increases in alkaline phosphatase, alanine aminotransferase and aspartate aminotransferase levels at 24 hours postdose. The erythrocyte-to-hepatocyte volume ratio in the livers of APAP-treated mice was also reduced by anti-Cd41 IgG pretreatment, suggesting a reduction in congestion or hemorrhage. The early (30 min) APAP-mediated consumption of glutathione (GSH) was unaffected by platelet depletion, suggesting that the reduction in injury was not a consequence of diminished APAP bioactivation. Activated platelets provide a thrombogenic surface for amplification of thrombin generation; indeed, platelet depletion significantly reduced the increase in plasma concentration of thrombin-antithrombin complexes, a biomarker of thrombin generation, at 3h. Addition of thrombin to cultures of primary hepatocytes in vitro did not affect APAP-induced cytotoxicity, suggesting that thrombin does not contribute to toxicity through a direct action on hepatocytes. Taken together, the results suggest that platelets support thrombin generation and contribute to congestion or hemorrhage and hepatocellular injury after APAP overdose. (Supported by NIH grant R01 DK087866.)

Purinergic Receptor Antagonist A438079 Protects against Acetaminophen-Induced Liver Injury by Inhibiting P450 Isoenzymes Not Inflammamome Activation.

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Platelet Depletion Reduces Acetaminophen Hepatotoxicity in Mice.

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Acetaminophen (APAP) overdose is the major cause of acute liver failure in the U.S. Previous studies identified decreases in blood platelet concentration in patients with APAP toxicity, but the role of platelets in APAP-induced liver injury is unclear. We tested the hypothesis that platelets contribute to liver injury in a mouse model of APAP hepatotoxicity. Blood platelet concentration was reduced 6-36 h after administration of APAP (300 mg/kg, ip) to mice, and immunofluorescence labeling showed increased platelets within livers at 2-6 h. Pretreatment of mice with anti-Cd41 IgG reduced platelet concentrations to 1/10th of control reduced liver injury at 6, 12 and 24 h after APAP administration, as evaluated by plasma alanine aminotransferase activity and the area of hepatic necrosis. The erythrocyte-to-hepatocyte volume ratio in the livers of APAP-treated mice was also reduced by anti-Cd41 IgG pretreatment, suggesting a reduction in congestion or hemorrhage. The early (30 min) APAP-mediated consumption of glutathione (GSH) was unaffected by platelet depletion, suggesting that the reduction in injury was not a consequence of diminished APAP bioactivation. Activated platelets provide a thrombogenic surface for amplification of thrombin generation; indeed, platelet depletion significantly reduced the increase in plasma concentration of thrombin-antithrombin complexes, a biomarker of thrombin generation, at 3h. Addition of thrombin to cultures of primary hepatocytes in vitro did not affect APAP-induced cytotoxicity, suggesting that thrombin does not contribute to toxicity through a direct action on hepatocytes. Taken together, the results suggest that platelets support thrombin generation and contribute to congestion or hemorrhage and hepatocellular injury after APAP overdose. (Supported by NIH grant R01 DK087866.)

The Role of Choline Depletion in Perfluorooctanesulfonate-Induced Hepatic Steatosis.

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In toxicological studies, perfluorooctanesulfonate (PFOS) exposure has resulted in hepatic steatosis and hypolipidemia, believed to be the result of decreased production and secretion of VLDL and HDL and increased uptake of VLDL-triglyceride. The hypothesis that PFOS produces hepatic steatosis via ionic sequestration of available choline necessary for the production of VLDL. Plausibility was supported by identification of an ion-triad between two molecules of choline and one molecule of PFOS in vitro. We sed male C57BL/6 mice either a control diet or a marginal methionine/choline-deficient (mMCD) diet, both with and without PFOS. After a two-week run-in period on diets without PFOS, control or mMCD diets containing 0, 30, 60, or 120 mg K+PFOS/kg diet were fed for three weeks. There was a dose-dependent increase in the relative liver weight in both control and mMCD fed mice. Diabetic PFOS was also associated with dose-dependent decreases in body weight, increases in hepatic triglyceride concentration, and increases in serum ALT, ALP, and bile acids, all with larger effects observed in mice fed mMCD compared to those fed the control diet. Serum and liver concentrations of PFOS were increased in a dose-dependent manner on both diets; however, serum PFOS concentrations were higher and liver concentrations lower in mMCD-fed mice compared to corresponding control-fed mice. This is surprising because the evidence suggested that PFOS-induced hepatotoxicity was exacerbated in the mMCD diet-fed mice. Metabolomic analysis demonstrated that PFOS caused a significant decrease in the hepatic concentration of many phosphatidylcholines in the PFOS-fed mice compared to controls. Further, the average serum concentration of choline was reduced by dietary PFOS. These studies are the first to provide evidence that PFOS may cause hepatic steatosis through depletion of choline required for hepatic VLDL production and export.

Identifying Prepgenerative Signaling after Acetaminophen-Induced Acute Liver Failure in Mice Using Incremental Dose Model.

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Overdose of acetaminophen (APAP) is the major cause of acute liver failure (ALF) in the US with very limited treatment options. Recent studies suggest that liver regeneration is a critical determinant of overall survival following APAP overdose and...
has highlighted the potential of regenerative therapies for ALF. However, the mechan- 
imism of liver regeneration following APAP-induced ALF is not completely clear. In the present study, we studied the major signaling pathways involved in liver re-
generation following APAP-induced ALF using an incremental APAP dose model. Two-month-old male C57BL/6 mice were treated with either 300 mg/kg (APAP300) or 600 mg/kg (APAP600) APAP and liver injury and regeneration was studied over a time course of 0 to 48 hr. Mice treated with APAP300 developed ex-
tensive liver injury followed by significant liver regeneration resulting in resolution of APAP-induced injury by 48 hr. In contrast, mice treated with APAP600 exhib-
ted significant injury but substantial decrease in liver regeneration. The inhibition of liver regeneration in APAP600 group was associated with decreased Cyclin D1 RNA and protein expression, decreased phosphorylation of Rh protein and sus-
tained expression of p21. Further analysis of several signaling pathways revealed dif-
f erences in pro-mitogenic signaling between APAP300 and APAP600. EGFR, C-
Met, and downstream activation of MAPK pathways were highly activated in APAP600, where regeneration was inhibited. However, canonical Wnt/b-catenin pathway and NF-kB signaling were activated in APAP300 where liver regeneration was stimu-
lated and inhibited in APAP600 group where regeneration was decreased. Finally, a TaqMan-based PCR array of 15 growth factor genes revealed rapid induc-
tion of several growth factors including HGF, VEGF, Kit-L, and PDGFβ specifically in APAP300 group. Taken together, our study has identified Wnt and NF-kB signaling as potential pathways that regulate liver regeneration after APAP over-
dose.

**635 Potential Hepatoprotective Effects of Licorice Root (Radix glycyrrhiza) Extract against Carbon Tetrachloride-Induced Hepatotoxicity in Isolated Rat Hepatocytes.**

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Radix glycyrrhiza is one of the native Mediterranean plants. Licorice root is a pop-
ular soft drink in Egypt. Literature cited the therapeutic effects of licorice. The present work is to evaluate the potential hepatoprotective effects of aqueous licorice root extract against the cytotoxic effects and the oxidative stress induced by carbon tetra-
chloride (CCL4) in isolated primary rat hepatocytes Hepatocytes were isolated by collagenase perfusion technique. Cytotoxicity was deter-
ded by assessing cell viability and leakage of cytosolic enzymes, such as lactate dehydrogenase (LDH), alanine aminotransferase (ALT) and aspartate aminotransferase (AST). Oxidative stress was assessed by determining reduced glutathione (GSH) level and lipid peroxidation as indicated by thioarbituric acid reactive sub-
stances (TBARS) production. Exposure of isolated rat hepatocytes to CCL4 (5mM) caused cytotoxicity and oxidative injury, manifested by loss of cell viability and sig-
nificant increase in LDH, ALT and AST leakage. As well as, CCL4 caused progres-
sive depletion of intracellular GSH content and significant enhancement of TBARS accumulation.

Preincubation of hepatocytes with either licorice (25 μM/mL) or silymarin (0.5 mM) which is a known hepatoprotective agent, ameliorated the hepatotoxicity and oxidative stress induced by CCL4, and induced significant improvement in cell viability, significant decrease in LDH, ALT and AST leakage, significant pre-
vention of GSH depletion and significant decrease in TBARS formation as com-
pared to CCL4 alone-treated cells. The present results indicate that CCL4 has a po-
tential cytotoxic effect in isolated rat hepatocytes; and licorice extract possess a highly promising hepatoprotective effects against CCL4 - induced hepatotoxicity a suppression of apoptosis. Following prolonged exposure, this pro-proliferative mi-
lieu along with suppression of apoptosis promotes the growth and progression of spontaneously transformed cells eventually producing hepatocellular adenoma.

The ability of CCZ and PPZ to elicit these key events in vivo was confirmed by measuring Cyp2b and Cyp3a induction as a surrogate for CAR activation and replicative DNA synthesis to assess hepatocellular proliferation. In addition, the requirement for CAR for increased hepatocellular proliferation was demonstrated for CCZ using CAR knock-out mice. Furthermore, reporter gene as-
says demonstrated that PPZ is a direct activator of both mouse (pottent) and human (weak) CAR. Following experimental demonstration of the proposed MOA for mice, the lack of human relevance was assessed by comparing the effects of PPZ or 
CCZ on primary cultures of mouse and human hepatocytes. Consistent with the in 
vivo observations, treatment of mouse hepatocytes with CCZ or PPZ resulted in in-
creases in Cyp2b, Cyp3a and replicative DNA synthesis. In contrast, treatment of human hepatocytes with these compounds resulted in increased Cyp2b and Cyp3a, but not replicative DNA synthesis. These data demonstrate a qualitative species dif-
f erence in response to CAR activation. The lack of proliferative response in human hepatocytes means it is reasonable to conclude that neither CCZ nor PPZ are he-
patotumourogogenic in humans.

**636 Eosinophils Not Neutrophils Mediate Halothane-Induced Liver Injury in Mice.**

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Liver eosinophilia has often been associated with drug induced liver injury, though its role in the etiology of this disease remains unclear. We decided to investigate this problem in a murine model of halothane-induced liver injury, where neutrophils have been reported to play a pathogenic role. When female Balb/c mice were ad-
ministered halothane, eosinophils were detected by flow cytometry in the liver within 12 hours and increased thereafter proportionally to liver damage. Infiltrating neutrophils did not increase significantly until 18 hours after halothane administra-
tion. Chemokines CCL11 and CCL24, which are known to attract eosinophils, in-
creased in response to halothane-treatment. Immunohistochemical staining for the 
cytotoxic eosinophil granule protein, major basic protein, revealed that eosinophils accumulated exclusively around areas of hepatocellular necrosis. The severity of HILL was decreased significantly when the study was repeated in wild-type mice partially depleted of eosinophils by pretreatment with Siglec-F antibody, while the
number of hepatic neutrophils remained unchanged. Conversely, selective deple-
tion of neutrophils by pretreatment with low concentrations of Gr-1 antible failed to reduce the extent of HILI when levels of eosinophils remained unchanged. The pathologic role of eosinophils was confirmed when halothane induced hepato-
toxicity was significantly reduced in the eosinophil lineage ablated AdhGata^{	ext{Cre}}^- mice, which are neutrophil competent. Our findings indicate that eosinophils, not neutrophils, have a pathologic role in HILI in mice and suggest that they may con-
tribute similarly in many clinical cases of DILI.

637 Effect of Ligand Activation of PPARβ/δ in Kupffer Cells. 
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Peroxisome proliferator-activated receptor β/δ (PPARβ/δ) can inhibit pro-inflammatory activities in the liver. Since activated Kupffer cells can modulate hepatic inflammation, the role of PPARβ/δ in modulating Kupffer cell activities was examined. Kupffer cells were isolated from wild-type, Pparβ/δ-null or Pparβ/δ-null mice expressing a DNA binding mutant form of PPARβ/δ in the Kupffer cells (PPARβ/δ DBM). Cultured Kupffer cells from the three genotypes phagocytized latex beads demonstrating relative purity. Ligand activation of PPARβ/δ in Kupffer cells increased expression of the PPARβ/δ target gene, adipocyte differentiation-related protein (Adip) in wild-type but not in Pparβ/δ-null or PPARβ/δ DBM Kupffer cells. Ligand activation of PPARβ/δ attenuated lipopolysaccharide (LPS)-induced expression of the pro-inflammatory gene, tumor necrosis factor-alpha (Tnfa) mRNA in Kupffer cell cultures from wild-type and PPARβ/δ DBM mice but not from Pparβ/δ-null mice. Wild-type Kupffer cells treated with LPS to in-
duce an M1 phenotype exhibited increased expression of the pro-inflammatory macrophage markers Tnft, interleukin-6 (Il-6), interleukin-1β (Il-1β) and chemokine ligand-4 (Ccl4) and ligand activation of PPARβ/δ attenuated these re-
sponses. Wild-type Kupffer cells treated with IL-4 to induce an M2 phenotype ex-
hibited increased expression of the anti-inflammatory markers arginase type-1 (Arg-1), macrophage galactose N-acetyl-galactosamine specific lectin-1 (Mgl-1), c-
type lectin domain family 7, member A (Clec7A) and interleukin-10 (Il-10), but ligand activation of PPARβ/δ did not influence these responses. Combined, results from these studies suggest that PPARβ/δ inhibits hepatic inflammation, at least in part, via transrepression of pro-inflammatory signaling in Kupffer cells. Since this effect can be found with a DNA binding mutant form of PPARβ/δ, this suggests that the attenuation of pro-inflammatory signaling could be due to direct protein-protein interaction of PPARβ/δ with other inflammatory signaling molecules such as NF-κB.

638 Hepatocyte Tissue Factor Triggers the Procoagulant Response Associated with Acetaminophen-Induced Liver Injury and Hepatocyte Transplantation. 
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The localization and regulation of tissue factor (TF) activity in hepatocytes (HPCs) is poorly understood and a role for HPC TF in vivo has not been established. We characterized the expression of TF by mouse HPCs and evaluated the role of HPC TF in models of HPC transplantation and acetaminophen (APAP) over-
dose. TF mRNA, protein, and total procoagulant activity (PCA) were significantly reduced in isolated HPCs and in liver homogenates from TF^{	ext{hetero}}^-/^-albumin-Cre mice (HPC^{	ext{Cre}}^- mice) compared to Tflox/flox mice (control mice), TF protein on the surface of intact HPCs had PCA that was reduced by cell-impermeable lysine conjugating reagents, but was unaffected by an inhibitory TF antibody that com-
petes for factor VII binding. Intact mouse HPCs clotted factor VII-deficient human plasma and TF-dependent factor Xa generation by HPCs occurred without exogenous factor VIIa. Thrombin generation in a model of HPC transplantation was dependent on donor HPC TF expression. Thrombin generation was also dra-
maically reduced in APAP-treated HPC^{	ext{Cre}}^- mice compared to APAP-treated con-
tral mice. The results indicate that TF expressed on the surface of mouse HPCs is preloaded with VII/VIIa, and that HPC TF is essential for coagulation induced by hepato
cellular injury in vivo. Moreover, the expression of TF by HPCs implies that most hepatotoxic responses are likely accompanied by activation of the extrinsic pathway of blood coagulation.

639 Hepatoprotective Effects of Fibrinogen(ogen) in Chronic Xenobiotic-Induced Cholestatic Liver Injury: Potential Involvement of Platelets. 
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Coagulation cascade activation and hepatic fibrin(ogen) deposition are evident in a model of chronic alpha-naphthylisothiocyanate (ANIT)-induced cholestatic liver injury. We have shown previously that complete fibrin(ogen) deficiency reduces liver necrosis in mice fed a diet containing ANIT. The mechanism whereby fib-
rin(ogen) deficiency worsens injury in this model is not known, nor is it clear whether stabilizing hepatic fibrin would be hepatoprotective. One possibility is that fibrin(ogen)-dependent platelet activation protects the liver from chronic ANIT diet-induced injury. We tested the hypothesis that stabilizing hepatic fibrin(ogen) inhibits ANIT-induced chronic cholestatic liver injury by supporting hepatic platelet accumulation/activation. Mice fed a diet containing 0.025% ANIT for 2 weeks developed liver injury consisting of multifocal acute hepatic necrosis and inflammation, periportal fibrosis, lymphocytic inflammation and bile duct ep
thelial hyperplasia. Immunofluorescent CD41 (alphad integrin) staining revealed a marked increase in platelets in livers of mice fed the ANIT diet compared to mice fed control diet. Administration of the anti-fibrinolytic drug tranexamic acid (1200 mg/kg, ip, bid) significantly reduced liver injury in mice fed the ANIT diet, but did not affect platelet accumulation. Administration of the platelet inhibitor clopidogel significantly increased serum alanine aminotransferase (ALT) levels and liver necrosis in mice fed the ANIT diet. Moreover, serum ALT activity was increased in mice deficient in protease activated receptor-4 (PAR-4), a primary thrombin recep-
tor on mouse platelets. The results indicate that stabilization of hepatic fibrin with antifibrinolytic therapy significantly reduces liver injury in mice fed ANIT diet. Moreover, results suggest that platelets are also hepatoprotective in mice fed the ANIT diet.

640 Effects of Acetaminophen and Oxidative Stress on Primary Human Hepatocytes Derived from a Steatotic Liver. 
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Primary human hepatocytes are a favored in vitro system for characterization of some types of hepatotoxicity. Donor demographics, disease states, and lifestyles re-
sult in significant donor to donor variability and can have a strong influence on the sensitivity of primary hepatocytes to hepatotoxins. Here, we describe a unique primary human hepatocyte system originating from the liver of a female donor (SMK) with severe metabolic syndrome and pre-conditioned steatosis. The cells had a lipid-rich phenotype that was clearly visible both using standard light mi-
croscopy and lipid fluorescent stain, which showed a 400% increase in signal inten-
sity compared to a standard lot of hepatocytes. In general, steatotic cells have en-
hanced sensitivity to the presence of reactive oxidative stress (ROS). To test whether SMK cells were more susceptible to ROS, SMK and cells from a normal donor were exposed to acetaminophen, a drug with a well characterized ROS-mediated mecha
nism of toxicity, at a dose range of 0.5 to 20 mM for 72h using standard culture conditions. SMK hepatocytes were 2.7-fold more susceptible to APAP induced cy-
totoxicity compared to their normal counterparts. The lipid signal was slightly re-
duced upon high levels of APAP. Total RNA was isolated from SMK both from na
ive and APAP treated cells and subjected to whole genome transcriptomic profil-
ing. When compared to a pool of non-diseased hepatocyte donors, naive SMK cells showed extensive gene expression changes that were largely associated with bioener
getic pathways with marked downregulation of lipid processing control and synthe
sis genes (e.g. SREBP). Agglomerative hierarchical cluster analysis revealed a clear distin
ction between SMK and its normal counterparts of the global expression pro-
files induced after treatment with APAP. In summary, this study demonstrates that hepatocytes with marked steatosis can serve as interesting molecular models for ROS-mediated hepatotoxic injury.
**641** Mesito-Dihydroguaiaretic Acid Inhibits Alcoholic and Nonalcoholic Fatty Liver by Antagonizing LXRα Activity.

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Collaborative regulation of liver X receptor (LXR) and steroid regulatory element binding protein (SREBP)-1 are main determinants in hepatic steatosis. Recent studies indicate that selective intervention of overlying functional LXRα in the liver shows promise in treatment of fatty liver. In the present study, we evaluated the effects of mesito-dihydroguaiaretic acid (MDGA) on LXRα activation and its ability to reverse fatty liver in mice. An LXRα co-activator recruitment assay and molecular docking analysis were performed to evaluate the binding of MDGA to the ligand-binding domain (LBD) of LXRα. The ability of MDGA to inhibit LXRα-dependent steatosis was investigated in an ethanol- or high-fat-diet (HFD)-induced steatosis model. MDGA inhibited activation of the LXRα-RBD by competitively binding to the site of agonist T0901317 and decreased the luciferase activity in LXRE-tk-Luc-transfected cells. MDGA attenuated expression of LXRE-dependent genes, and the decreased liver weights in CARKO mice were compared with control mice. MDGA reduced the expression of LXRα co-activator protein RIP140 in HFD-fed mice. While MDGA decreased the expression of LXRα, SREBP-1, SCD-1, and FAS proteins, genes associated with reverse cholesterol transport such as ABCA1 and ABCG1 were not affected. These results demonstrate that MDGA has the potential to reverse alcoholic and nonalcoholic steatosis mediated by selective inhibition of LXRα in the liver.

**642** Toxicity of Diethylnitrosamine (DEN) in the Liver of Constitutive Andorostane Receptor (CAR)-Knockout (KO) Mice.

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CARKO mice showed high mortality after injection of DEN. To clarify the cause of high susceptibility in CARKO mice to the toxicity of DEN, 6-week-old male CARKO and C3H mice were intraperitoneally injected 90 mg/kg body weight of DEN, and their livers were collected at days 1, 3, 7 and 14 after DEN treatment. Relative weight, pathological changes, mRNA expression and enzyme activity levels of CYP2B10 and CYP2E1 in the liver of CARKO mice were compared with controls injected saline or with wild mice. DEN significantly reduced relative liver weights in both genotypes and the decreased liver weights in CARKO mice were significantly lower than wild mice at days 7 and 14. Microscopically, pale cytoplasma of the hepatocytes in the centrilobular area was observed by DEN treatment at day 1, and the area of this change was larger in CARKO than in wild mice. DEN treatment also increased slight to mild single necrosis of hepatocytes and mononuclear cell infiltration in the centrilobular area from day 1 and from day 3, respectively. The intensities of single necrosis and mononuclear cell infiltration in CARKO mice were comparable to wild mice. While the expression level of Cyp2b10 in the liver of wild mice increased by DEN with or without significance, compared with their controls, the level in CARKO mice remarkably diminished in both control and DEN groups at all time points. As for testosterone hydroxylase activity (CYP2B10), there were no difference between controls and DEN groups or both genotypes at days 1 and 7. DEN treatment significantly decreased Cyp2e1 and aniline hydroxylase activity (CYP2E1) in both genotypes at day 1, compared with each control, and the decreased level of aniline hydroxylase activity was significantly lower in CARKO than in wild mice. Judging from these results, high susceptibility in CARKO mice to the toxicity of DEN might be caused by immediate decrease of CYP2E1 activity after DEN treatment.

**643** Leptin Modulates Toxicity Associated Steatohepatitis through Peroxynitrite in Kupffer Cells.

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Progression from steatosis to toxin-mediated steatohepatitis (TASH) lesions is hypothesized to require a second hit. These lesions have been associated with increased oxidative stress, often ascribed to high levels of leptin and other proinflammatory mediators. Here we have examined the role of leptin in inducing oxidative stress and Kupffer cell activation in toxin-mediated steatohepatitic lesions of obese mice. Male C57BL/6 mice fed with a high fat diet (60%kcal) at 16 weeks were administered CCl4 to induce steatohepatitic lesions. Approaches included use of immunosuppression for measuring free radical stress, gene-deficient mice for leptin, p47 phox, iNOS and adoptive transfer of leptin primed macrophages in vivo. Diet-induced obese (DIO) mice were treated with CCl4, mice primed with VC and leptin receptor expression. Oxidative stress was significantly elevated in DIO mouse liver but not in OB/OB mice, or in DIO mice treated with leptin antibody. In OB/OB mice, leptin supplementation restored markers of free radical generation. Markers of free radical formation were significantly decreased by the peroxynitrite decomposition catalyst FeTPPS, the iNOS inhibitor 1400W, the NADPH oxidase inhibitor apocynin, or in iNOS or p47phox-deficient mice. These results correlated with the decreased expression of TNF-alpha and MCP-1 and decreased leptin receptor expression. Kupffer cell depletion eliminated oxidative stress and inflammation, whereas in macrophage-depleted mice, the adoptive transfer of leptin-primed macrophages significantly restored inflammation. These results, for the first time, suggest that leptin action in macrophages of steatotic liver through induction of iNOS and NADPH oxidase caused peroxynitrite-mediated oxidative stress thus activating Kupffer cells and modulating leptin receptor expression in the liver. (NIH-R00ES19875)

**644** Occupational Vinyl Chloride Exposures Are Associated With Significant Changes to the Plasma Metabolome: Implications for Toxicant Associated Steatohepatitis.

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Occupational vinyl chloride (VC) exposure has been associated with steatohepatitis (TASH) and liver cancer, although the modes of action are unknown. Metabolomics has recently been utilized for the evaluation of drug-induced liver injury, but has not previously been performed for occupational VC exposures. Plasma samples from 17 highly-exposed VC workers without liver cancer and 27 unexposed healthy volunteers were obtained from a specimen bank. GC/MS (Thermo-Finnigan Trace-DQG fast-scanning single-quadrupole mass spectrometer) and LC/MS2 (Waters ACQUITY UPLC, Thermo-Finnigan LTQ mass spectrometer) were performed. Software was used to match ions to a library of standards for metabolite identification and quantitation by peak area integration. Statistical significance was determined using Welch’s t-tests. 613 unique named metabolites were identified. Of these, 189 metabolites were significantly increased in the VC exposure group while 94 metabolites were significantly decreased. The most striking differences occurred in lipid metabolites. Essential (7 of 7) and long chain free fatty acids (19 of 19, including arachidonic acid, 6 fold) were significantly increased with VC exposure. Lipid peroxidation products were likewise increased by VC exposure: monoxygenated fatty acids (8 of 9, including 5-HODE, 211 fold); fatty acid dicarboxylates (8 of 13); oxidized arachidonic acid products (7 of 7, including 5-HETE, 9-HETE, and 15-HETE, up to 616 fold). Other arachidonic acid products including leukotriene B4 (52 fold) were also up-regulated with VC exposure. Abnormalities were also noted in amino acid metabolism, and particularly the transmethylation and transsulfuration pathways. VC exposure was associated with increased plasma free fatty acids and lipid peroxidation products. Lipotoxicity, pro-inflammatory lipid peroxidation products, and impaired transmethylation/transsulfuration pathways represent novel modes of action for VC hepatotoxicity.

**645** Pregnancy-Related Lactogenic Hormones Alter the Expression of Uptake and Efflux Transporters in Primary Human Hepatocytes.

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The secretion of lactogenic hormones prolactin (PRL) and placental lactogen (PL) increases steadily throughout pregnancy and reaches the highest levels just prior to parturition. The effects of lactogenic hormones on the expression of hepatic transporters critical for the excretion of endogenous chemicals, drugs, and toxicants have not been characterized. To investigate the regulation of transporters by lactogenic hormones, sandwich-cultured, freshly isolated human hepatocytes from three adult female donors were treated for 72 hours with vehicle (carbonate-buffered saline),
human PRL (150 ng/ml) or human PL (6 μg/ml), and total RNA was isolated. Gene expression was quantitated by microarray analysis and validated with quantitative real-time PCR (qPCR). qPCR analysis shows PRL and PL have differential effects on the gene expression of hepatic transporters. PRL caused the up-regulation of mRNA for the uptake transporter OAT2 by 60%, and down-regulation of the efflux transporters MRP3, MRP6, and BCRP between 15 and 20%. PL decreased mRNA expression for all uptake transporters tested, significantly for NTCP, OCT1, OATP1B1, and OATP1B3. In addition, PL increased efflux transporter gene expression for ABCA1 and BSEP by 100%, but reduced MDR1, MRP2, MRP3, MRP5, and BCRP mRNAs. Taken together, these data suggest hormones secreted during pregnancy may globally suppress the maternal gene expression of many uptake and efflux transporters crucial for chemical excretion by the liver. Supported by ES020522, DK080774, ES07148, ES050522, HD065532.

464 Acrolein Cytotoxicity in Hepatocytes Involves Endoplasmic Reticulum Stress, Mitochondrial Dysfunction, and Oxidative Stress.

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Acrolein is a common environmental, food and water pollutant and a major component of cigarette smoke. Also, it is produced endogenously via lipid peroxidation and cellular metabolism of certain amino acids and drugs. Acrolein is cytotoxic to many cell types including hepatocytes; however the mechanisms are not fully understood. We examined the molecular mechanisms underlying the acrolein cytotoxicity in primary human hepatocytes and hepatoma cells. Acrolein, at pathophysiological concentrations, caused a dose-dependent loss of viability of hepatocytes. The death was apoptotic at moderate and necrotic at high concentrations of acrolein. Acrolein exposure rapidly and dramatically decreased intracellular glutathione and overall antioxidant capacity, and activated the stress-signaling MAP kinases JNK, p42/44 and p38. Our data demonstrate for the first time in human hepatocytes, that acrolein triggered endoplasmic reticulum (ER) stress and activated eIF2α, ATF-3 and -4, and Gadd153/CHOP, resulting in cell death. Notably, the protective/adaptive component of ER stress was not activated, and acrolein failed to up-regulate the protective ER-chaperones, GRP78 and GRP94. Additionally, exposure to acrolein disrupted mitochondrial integrity/function, and led to the release of pro-apoptotic proteins and ATP depletion. Acrolein-induced cell death was attenuated by N-acetyl cysteine, phenyl-butyric acid, and caspase and JNK inhibitors. Our data demonstrate that exposure to acrolein induces a variety of stress responses in hepatocytes, including GSH depletion, oxidative stress, mitochondrial dysfunction and ER stress (without ER- protective responses) which together contribute to acrolein toxicity. Our study defines basic mechanisms underlying liver injury caused by reactive aldehyde pollutants such as acrolein.

467 Interspecies Differences in Progression from Simple Liver Steatosis to Fibrosis Are Associated with Altered Hepatic Iron Metabolism in Mice.

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Nonalcoholic fatty liver disease (NAFLD) is a major health problem and a leading cause of chronic liver disease in the United States and other developed countries. The pathogenesis of NAFLD is often conceptualized as a two-step process that consists of hepatic triglyceride accumulation leading to steatosis as a first step, followed by a second step that includes oxidative stress, inflammation, and fibrogenesis. However, there is a lack of consensus on the significance of these events and the underlying mechanisms responsible for the disease progression and, more importantly, for differences in inter-individual disease severity and sensitivity. In a previous study we demonstrated that the inter-strain variability in severity of NAFLD was associated with down-regulation of the lipid metabolism first step (hypo-glycosylation). This study further supports the hypothesis that hypo-glycosylation in the first step of lipid metabolism is a key determinant of the interstrain variability in severity of NAFLD and its further progression to fibrosis, which was characterized by hepatic stellate cell activation and accumulation of fibrous markers, was associated with the extent of hepatic iron metabolism deregulation. This was evidenced by strain-dependent alterations in the expression of iron- regulatory genes (Tfrc, Fth1, Slc40a1, and Hfe2), which was correlated tightly with the degree of hepatic fibrotic changes in the livers of the mice fed the CFD diet.

468 Dose-Dependent Hepatic Physiological, Histopathological, and Gene Expression Responses in C57BL/6 Mice following Repeated TCDD Exposure.

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2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD) is a potent aryl hydrocarbon receptor (AhR) agonist. We have previously reported that exposure to a single oral dose of 1 μg/kg TCDD induces hepatic steatosis in mice. A complementary 28 day repeated-dose dose-response study was performed to further investigate the effects of TCDD on steatosis and its progression to steatohepatitis and cirrhosis. C57BL/6 mice were gavaged every 4 days with sesame oil vehicle control or 0.001 - 30 μg/kg TCDD. Although repeated dosing did not significantly affect body weight gain, gonadal white adipose tissue (gWAT) weight was 50% lower at 30 μg/kg while relative liver weight was increased at 3, 10, and 30 μg/kg TCDD. Histopathology revealed dose dependent alterations with minimal centricrinar microvesicular lipid accumulation (hepatic steatosis) at 3 μg/kg, mild to moderate hepatic lipid accumulation with mild inflamation at 10 μg/kg (steatohepatitis), and widespread micro- and macro-vesicular lipid accumulation in the centricrinar, mid-zonal, and periporal regions of the liver, inflammation, and perivenular fibrosis suggesting early cirrhosis at 30 μg/kg. QRTPCR revealed dose-dependent induction of Cyp1a1, Cyp1a2, Cyp1b1, Dnaj, Gsp, Gap2, Notech1, Nqat1, Plaq212a, Serpinbda, Sre1, Tiparp, and Trafip2, as well as down-regulation of G61 and Pek1. Dose-response modeling (ToxpResponse modeler) revealed similar ED50’s to 24h single dose TCDD studies. These results suggest that simple steatosis can progress to steatohepatitis with fibrosis following repeated TCDD exposure, due to persistent AhR-mediated differential gene expression associated with lipid transport, processing, and metabolism. Funded by SPP P42ES04911.

469 Impaired Glycosylation and Membrane Localization of Uptake and Efflux Transporters in Human Nonalcoholic Fatty Liver Disease.

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The prevalence of non-alcoholic fatty liver disease (NAFLD) and the more severe non-alcoholic steatohepatitis (NASH) is estimated to be 30-40% and 5-17%, respectively. We have previously demonstrated a loss of N-linked glycosylation and decreased membrane localization of the efflux transporter ABC2C2 in rodent and human NASH resulting in the altered disposition of drugs. N-linked glycosylation of proteins has been shown to be critical for proper protein folding and trafficking to the plasma membrane. The purpose of this study was to assess the transcriptomic expression of genes involved in protein glycosylation and processing through the endoplasmic reticulum (ER), and to determine the effect of altered glycosylation on key drug transporters during the progression of human NAFLD. For this study, human liver samples diagnosed as healthy, steatosis, NASH with fat, and NASH without fat were analyzed. Using bioinformatic methods we discovered that genes involved in protein processing in the ER and biosynthesis of N-glycans were significantly enriched for downregulation in NAFLD progression. Included in the N-glycan biosynthesis category were genes involved in the oligosaccharyltransferase complex, N-glycan quality control, N-glycan precursor biosynthesis, N-glycan trimming to the core, and N-glycan extension from the core. In contrast to down-regulation of these genes, N-glycan degradation genes were unaltered in the progression to NASH. Immunoblot analysis of the uptake transporters OATP1B1 and OATP1B3 and the efflux transporters ABC2C2 and BCRP demonstrated a significant loss of glycosylation, while immunohistochemical analysis revealed impaired membrane localization of ABC2C2. We propose that the loss of glycosylation and impaired membrane localization of key uptake and efflux transporters in human NASH is a potential mechanism for the occurrence of altered drug disposition in NASH.
High Dietary Fructose Induced Copper Deficiency Contributes to Alcoholic Liver Disease Progression.

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Background/Aims: The increased consumption of fructose parallels the increased prevalence of obesity and the metabolic syndrome in the United States and worldwide. Noteworthy is that obesity potentiates the severity of alcohol-induced liver injury. High fructose feeding impairs intestinal copper absorption and leads to copper deficiency in rodents. Moreover, copper deficiency is associated with the decreased antioxidant defenses and mitochondrial dysfunction. In this study, we investigated whether high fructose diet induced obesity potentiates chronic alcohol drinking induced liver injury.

Methods: Six-week-old male C57BL/6 mice were fed either control diet or high fructose diet (45%) for 16 weeks. 20% (v/v) ethanol was given ad libitum after 8-weeks high fructose feeding until the end of the experiment. Copper status, hepatic injury and steatosis were assessed.

Results: High fructose diet feeding led to copper deficiency as indicated by decreased plasma ceruloplasmin, Cu, Zinc superoxide dismutase (SOD1) and increased copper chaperone for SOD1 (CCS). Liver injury was significantly induced in mice fed with high fructose diet followed by chronic alcohol drinking as evidenced by robust increased plasma ALT, AST, chemokine (mouse KC) and liver histology. Both liver triglyceride and plasma triglyceride were significantly increased with chronic alcohol drinking, however, neither additive nor synergistic effects was observed in mice fed with high fructose diet followed by chronic alcohol drinking.

Conclusions: Our data suggest that high fructose diet-induced obesity may potenti- ate chronic alcohol drinking induced liver injury. High fructose feeding induced copper deficiency might be a priming factor to chronic alcohol drinking induced liver injury. The potential mechanism might relate to the decreased antioxidant defense caused by copper deficiency.

Effect of Diallyl Disulfide on the Cytokine Expression in Cadmium Chloride-Treated Liver Cells.


Cadmium is one of the most hazardous metals in the environment. Studies have shown that exposure to cadmium causes damage to many organs that may alter biologic- ological activities of the cells and may lead to cancer in mammalian systems. In the presence of toxicants, cells produce various cytokines in the body to reduce the toxic effect of the toxicants. Diallyl disulfide (DADS), an organosulfur compound in the garlic extract has been used in many countries as a preventive compound for various diseases. DADS can act as chelator and/or as an antioxidant. This project was designed to study the preventive effect of DADS on the cytokine expression in cadmium chloride (CdCl2) treated normal rat liver CRL-14439 cells. For the viability assay, the cells were treated with CdCl2 alone (0, 50 and 150 μM), DADS alone (150 and 300 μM) or co-treated with 2 h pre-treatment of DADS (150 and 300 μM) prior to CdCl2 (150 μM) for 24 h and the cells viability was measured by the crystal violet assay. For the cytokine array analysis, the cells were treated with CdCl2 alone (0, 50 and 150 μM), DADS (60%, 90% and 100%), and 2 h pre-treatment of DADS (150 μM) prior to CdCl2 (150 μM) for 6 h and the cytokines expression was measured using the Ray Biotech human cytokine array 7 kit. Viability results revealed that 150 μM of DADS showed the greatest protection against CdCl2 toxicity. In the cells treated with CdCl2 alone, 22 cytokines were up-regulated (fold change > 2) and 2 cytokines were down-regulated. Flow cytometry results revealed that the pattern of cytokines expression observed was reversed, indicating the protective effect of DADS against CdCl2 toxicity. Cytokines that were up-regulated in CdCl2 alone are involved in the alterations of cell cycle control system and cell mediated immunity. The present study clearly shows the protective effect of DADS on the cytokine expression in the CdCl2 treated liver cells and suggests that DADS can be used as preventative agent for cadmium toxicity.

The Effect of Fenugreek Leaf Extract on Gene Expression Profile of Cadmium Chloride Treated Normal Rat Liver Cells.


Rationale and Scope of the Study: Cadmium is a toxic and carcinogenic metal pollutant that has been known to cause DNA damage. It targets many human organs, mainly lungs, liver, and kidneys. Many chemo-preventive agents have been used against the toxic effect of many heavy metals. Fenugreek belongs to the family Leguminosae and well known for its medicinal value. In our study, the effect of Fenugreek Leaf Extract (FLE) on cytoprotective ability and toxicological profile in cadmium treated rat liver cells was evaluated. Experimental Procedure: The cells were treated with CdCl2 (0, 25 μM) alone or pretreated with FLE (0.005μg/ml) for 4 h followed by CdCl2 (25 μM) for 36 h or 48 h at 37°C in a 5% CO2 incubator. The viability was measured by crystal violet dye staining method. The total gene profiles were determined using RG230 PM whole genome microarray which was processed by Affymetrix Gene Atlas system. The Partek Express software analyzes the genes up or down regulated in the treated samples. The Partek pathway software identifies the pathways affected by the treatment. Results: In CdCl2 alone treated cells, the viability was reduced to 37.1%, while in the cells pretreated (4 h) with FLE followed by CdCl2, the viability was increased to 102% respectively, in comparison to the control cells (100%). In CdCl2 alone treated cells, out of 31,139 genes on the array 61 were up regulated (>2 fold) and 124 were down regulated (≤2 fold) respectively in comparison to control cells. In the cells pretreated with FLE followed by CdCl2, 181 genes were up regulated (>2 fold) and 161 genes were down regulated (≤2 fold). The main pathways affected in above treatment groups were ribosome, TCA cycle and DNA replication. Conclusion: Our results indicate that pretreatment with FLE for 4 h affects gene expression profile in the cadmium treated cells. The alteration in the genes expression in the FLE pretreated cells may be responsible for the protective effect against cadmium toxicity. Therefore, our study suggests that fenugreek leaves can be used to reduce cadmium toxicity.

Evidence for a Functional Keap1-Nrf2 Cell Defence Pathway in Human Liver.

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The transcription factor Nrf2 regulates a battery of cell defence processes that protect against drug-induced liver injury in rodents, yet very little is known regarding the role of Nrf2 in regulating resistance to drug-induced stress in man. Utilizing hepato- cytes isolated from surgically resected healthy liver tissue (n=6), we provide the first comprehensive delineation of the Nrf2 pathway in human liver. Hepatocytes were isolated using a standard collagenase perfusion protocol, seeded onto Type I collagen-coated culture plates and reverse-transfected with 20 nM of an siRNA duplex targeted against Nrf2, Keap1 or a scrambled non-targeting control siRNA duplex. siRNA depletion of Nrf2 was associated with a decrease in the mRNA and protein levels of the established Nrf2 target genes GCLC, NQO1 and SRXN1. Furthermore, following siRNA depletion of Keap1, the cytosolic repressor of Nrf2, the total cellular abundance of Nrf2 protein was increased, as was the mRNA and protein levels of GCLC, NQO1 and SRXN1. The potent Nrf2 activator CDDO-Me induced a robust stabilisation of Nrf2 in human hepatocytes, and this was associ- ated with a time and Nrf2-dependent increase in the mRNA and protein levels of GCLC, NQO1 and SRXN1. Under the same conditions, CDDO-Me provoked a
Mitochondrial function is very important in cellular homeostasis and keeping a proper energy supply for eukaryotic cells is essential in the fulfillment of the tissues energy-demand. The main objectives of this work concerned Dibenzofuran effects on mitochondrial function. We isolated mitochondria from rat liver and incubated them with Dibenzofuran to analyze the effects of this pollutant at the level of mitochondrial function. The effects of Dibenzofuran exposure include a markedly increase in the lag phase that follows depolarization induced by ADP, indicating an effect in the phosphorylative system. Experiments performed using carboxyatractyloside (CAT) suggested an interaction of Dibenzofuran with the ANT carrier. Dibenzofuran exposure also produces an inhibition of mitochondrial permeability transition and an increase in calcium retention capacity, which may also be explained by a putative interaction of Dibenzofuran with ANT.

Clarifying the role of pollutants in some mechanisms of toxicity, such as unbalance of bioenergetics status and mitochondrial function, may help to explain the progressive and chronic evolution of diseases derived from exposure to environmental pollutants.

Trovafloxacin (TVX) is a fluoroquinolone antibiotic associated with idiosyncratic drug-induced liver injury (IDILI) in humans. The mechanism underlying this toxicity is unknown. A animal model of IDILI in mice revealed that TVX synergizes with a concurrent inflammatory stress from bacterial lipopolysaccharide (LPS) to result in liver injury. This hepatotoxic interaction depended upon prolongation of the LPS-induced appearance of tumor necrosis factor-alpha (TNF) in the plasma of animals coexposed to TVX. We established a model in vitro in RAW 264.7 murine macrophages (RAW cells) in which exposure to TVX alone or in combination with LPS increases both cellular TNF mRNA and TNF protein released into the culture medium. TVX is a bacterial topoisomerase inhibitor, and in a cell-free assay it inhibited mammalian topoisomerase II-alpha as well. Interestingly, TVX also caused a concentration-dependent increase in DNA double-strand breaks (DSBs) in RAW cells. Inasmuch as DSBS can activate kinases such as ataxia telangiectasia mutated (ATM) and Rad3-related (ATR), kinase inhibitors were tested for their ability to diminish TVX-induced synthesis and release of TNF from RAW cells. A selective ATM inhibitor (KU55933) had no effect on TNF mRNA or TNF release from TVX-exposed RAW cells. In contrast, a novel ATR inhibitor (NU6027) significantly attenuated the TVX-induced increase in TNF mRNA. NU6027 also eliminated the interaction between TVX and LPS, significantly decreasing TNF release in cells treated with TVX-LPS to the level induced by LPS alone. These findings suggest that ATR plays an important role in TNF expression in response to TVX exposure. (Supported by NIH grant R01 DK061315 and T32 ES007255)
uptake. The phosphoinositide 3'-kinase (PI3K)/Akt signaling pathway plays an important role in glucose metabolism, in part by regulating the activity of forkhead box class O (FoxO) transcription factors. The present study aimed at investigating the effect of As³⁺ (arsenate) on FoxO activity in human hepatoma cells, the role of PI3K/Akt signaling therein and the modulation of the expression of FoxO target genes, such as that of selenoprotein P (SelP), a hepatocyte causing insulin resistance. We analyzed changes in phosphorylation of Akt and FOXO1α and β in HepG2 human hepatoma cells in response to As³⁺ by Western blotting and found a concentration-dependent increase in phosphorylation of Akt at S473, with arsenite being effective already at low micromolar concentrations. Arsenite exposure caused phosphorylation of FoxO1a and FoxO3a at sites known to be phosphorylated by Akt, T32 and T24, respectively. Phosphorylation of FoxOs was prevented by wortmannin, pointing to the involvement of PI3K. The functional inactivation of FoxOs by As³⁺ was observed in an ELISA-based DNA binding assay and at the level of FoxO target gene expression: SelP and glucose-6-phosphate dehydrogenase mRNA levels were clearly downregulated after 24h exposure to nanomolar As³⁺ concentrations, as demonstrated by real-time RT-PCR. Curiously, As³⁺ showed a biphasic effect on SelP protein levels, inducing a small increase in the nanomolar and a distinct decrease in the micromolar concentration range. In conclusion, As³⁺ may perturb cellular signaling pathways involved in fuel metabolism: it stimulates insulin-like signaling in HepG2 cells and affects the expression of FoxO target genes and the release of the hepatokine SelP, which is known to modulate insulin sensitivity.

**660 The Role of Multidrug Resistance Protein 4 (Mrp4, Abcc4) in Protecting against Acetaminophen (APAP)-Induced Hepatotoxicity.**

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Mrp4 is a member of the ABC family of membrane transport proteins that functions as an efflux transporter for a wide variety of substrates. Previous studies in our laboratory have shown that Mrp4 is the most significantly induced hepatic transporter by toxic acetaminophen (APAP) treatment. The basal expression of Mrp4 in normal human and naive C57BL/6J mouse liver is very low. However, Mrp4 expression is greatly induced by exposure to hepatotoxicants that cause oxidative stress, suggesting that this transporter is up-regulated to assist in defending against cellular stress. Additionally, our laboratory has demonstrated that a low hepatotoxic dose of APAP protects against subsequent administration of a higher dose of APAP. This phenomenon is referred to as APAP autoprotection and liver Mrp4 induction is even more pronounced in this mouse model. Despite these findings, the precise role that enhanced Mrp4 expression plays in protecting against APAP hepatotoxicity remains unclear. To study this, we analyzed the responsiveness of Mrp4 heterozygous mice (Mrp4+/−) to APAP hepatotoxicity and the ability of these mice to exhibit resistance to APAP upon toxicant re-exposure. The results showed that APAP hepatotoxicity in Mrp4+/− mice is significantly higher than in wild-type mice. However, the ability of these mice to develop tolerance to APAP liver injury is unchanged. Analysis of potential differences in gene expression of other hepatic membrane transporters, oxidative stress-related genes and drug detoxification enzymes revealed that there are no significant changes that could account for the differential susceptibility of Mrp4+/− mice to APAP toxicity. Based on these results, we conclude that Mrp4 plays a critical role in preventing APAP-induced hepatotoxicity but that its role in APAP autoprotection may be less substantial.

**661 Dual Role of Macrophages in Injury and Repair in Acetaminophen-Induced Hepatotoxicity.**

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Activated macrophages and the mediators that they release play a key role in acetaminophen (APAP)-induced hepatotoxicity; however, their role is dependent on their phenotype and timing of appearance in the liver. Whereas classically activated M1 macrophages contribute to tissue injury, alternatively activated M2 macrophages are involved in tissue repair. To assess the role of these macrophage subpopulations in APAP hepatotoxicity, we used gadolinium chloride (GdCl₃) and clodronate liposomes (CL), which block M1 and M2 macrophages, respectively. Male Long Evans hooded rats were treated with GdCl₃ (7 mg/kg, iv) or CL (0.4 ml/100 g body weight), 24 and 48 h prior to APAP (600 mg/kg, ip). Cells were isolated 24 h later. Treatment of rats with APAP resulted in a decrease in CD163+ resident repair macrophages in the liver after 24 h, with no significant effect on infiltrating CD11b+/CD68+ M2 repair macrophages. Pretreatment of rats with GdCl₃, which protects against APAP hepatotoxicity, resulted in a decrease in macrophage expression of inducible nitric oxide synthase, a prototypical marker of classical activation. This was associated with an increase in CD163+ resident and CD11b+/CD68+ infiltrating repair macrophages in the liver. Conversely, CL pretreatment increased APAP hepatotoxicity, markedly decreased both of these repair macrophage populations. Expression of markers of alternative M2 macrophage activation including STK, macrophage stimulating protein, and arginase-1, as well as the anti-inflammatory cytokine IL-10 were also reduced in CL treated rats. Taken together these data support the concept that macrophages play distinct role in APAP hepatotoxicity depending on their functional capacity. Supported by NIH GM034310, ES004738, CA132624, AR055073 and ES050222.

**662 Induction of Hepatic Bcrp Transporter Expression in Mice Treated with Perfluorooctanoic Acid.**


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Perfluorooctanoic acid (PFOA) is an industrial chemical that has been associated with negative health outcomes. The purpose of this study was to determine whether PFOA alters the expression of the breast cancer resistance protein (BCRP/ABCG2), an efflux transporter found in hepatocytes and renal proximal tubule cells, in mice and to determine whether PFOA interferes with BCRP transporter in vitro. To test this, Bcrp mRNA and protein expression were measured in the kidneys and livers of male C57BL/6J mice treated with PFOA (1 or 3 mg/kg/d po) for 7 days. In addition, ATPase and membrane vesicle experiments were used to assess whether PFOA alters human BCRP activity. As expected, PFOA treatment increased liver weights as well as the mRNA and protein expression of the known target, cytochrome P450 4A14, in the liver and kidneys of mice. Compared to vehicle-treated control mice, PFOA treatment increased hepatic, but not kidney, Bcrp mRNA and protein between 2- and 3-fold. Immunofluorescent staining revealed enhanced canicular Bcrp staining in liver sections from PFOA-treated mice. In the ATPase assay, PFOA decreased transporter activity in sulfasalazine-activated BCRP membranes. In addition, PFOA inhibited BCRP-mediated transport in membrane vesicles between 47 and 69% at high concentrations (>25 μM). In conclusion, PFOA increases hepatic Bcrp expression in mice and may inhibit human BCRP function at high concentrations. Supported by DK080774, ES020522, ES020721, ES005022 and ASPET SURF.

**663 Xenosensors Mediated Regulation of ATP Binding Cassette Transporter B6 (ABCB6).**

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Cytochromes P450 (P450s) are induced in response to therapeutic drugs and environmental contaminants, leading to increased detoxification and elimination of the xenobiotics. Xenosensors like PXR, CAR and AhR play a major role in drug and environmental contaminant induced expression of P450s. However, it is not known if exposure to xenobiotics induces ABCB6 expression, to contribute to heme biosynthesis similar to those of P450s. ABCB6, a member of the ABCB family of membrane transport proteins, is responsible for the secretion of heme into the bile ducts and plays a critical role in heme metabolism. ABCB6 is a xenobiotic transporter expressed in the liver, kidney, and intestine. Further, promoter activation studies demonstrate that both AhR and CAR induce the expression of Abcb6 in mouse liver. However, it is not known if exposure to xenobiotics induces ABCB6 expression, to contribute to heme biosynthesis. Our previous study demonstrated that xenobiotics induce ABCB6 expression, to contribute to heme biosynthesis. This study is the first to demonstrate direct transcriptional activation of both mouse and human ABCB6 by xenobiotics similar to those found in inducible P450s. The results presented in this work have both pharmacological and toxicological significance. AhR ligands have the ability to precipitate porphyria and both AhR and CAR ligands have been shown to promote cell growth and proliferation during cancerogenesis. Thus, ABCB6 induction by xenosensors AhR and CAR could be a potential contributing factor in drug induced porphyria and carcinogenesis.
664  Hepatic Flavin-Containing Monoxygenase-3 (FMO3) Protein Is Inducible by Acetaminophen Treatment.

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Flavin-containing monoxygenase 3 (FMO3) is a drug-metabolizing enzyme with functions similar to CYP450s. Hepatic expression of human FMO3 gene is highly variable and is important in the detoxification of xenobiotics. Fmo3 was thought to be non-inducible but recent data showed that activation of the Ah receptor induces Fmo3 mRNA in mice. Recent microarray work by our group also showed a drastic induction of liver Fmo3 mRNA expression in a mouse model of resistance to acetaminophen (APAP) hepatotoxicity (auto-protection). In addition to APAP, alpha-naphthyl isothiocyanate treatment and bile duct ligation in mice markedly increase Fmo3 gene expression. The purpose of this study was to evaluate the Fmo3 gene regulation and protein expression during APAP hepatotoxicity. Among all Fmo isoforms analyzed for mRNA expression, Fmo3 was most significantly induced following a single dose of APAP (~400mg/kg). Additionally, Fmo3 mRNA levels increased significantly by APAP, its protein expression was marginally changed. Consistent with this, the catalytic activity of Fmo3 measured by oxygenation of methimazole did not change significantly either. By contrast, both Fmo3 mRNA and protein expression are significantly higher in mice pretreated and re-exposed to APAP (auto-protection model). In agreement with greater Fmo3 protein expression in livers of auto-protected mice, its catalytic activity was also significantly higher. In summary, the dramatic changes in Fmo3 gene expression produced by a single dose APAP are not accompanied by concomitant changes in protein and enzyme function, whereas livers from mice pretreated and re-exposed to APAP exhibit induction of both Fmo3 protein expression and catalytic function. Additional work is currently underway to determine the functional significance of enhanced Fmo3 protein function and which factors and signaling pathways mediate Fmo3 gene and protein expression during APAP toxicity. Taken together, these findings establish for the first time induction of not only Fmo3 gene expression, but also significant protein levels and function. Supported by NIH DK090557.

665  Vanin-1 Knockout Mice Exhibit Alterations in Compensatory Immune Infiltration and Hepatocyte Proliferation following Acetaminophen Toxicity.

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Previously we have shown that Vanin-1 (Vnn1) knockout mice are more susceptible to APAP hepatotoxicity (400mg/kg, i.p.) despite no differences in hepatic glutathione (GSH) content or gene expression of APAP metabolizing enzymes or transporters. Here we show that in vitro, livers from both genotypes showed similar capacities to bioactivate APAP to its reactive metabolite (~1.8 nmol APAP-Gluc/min/mg protein) and to detoxify the parent compound by glucuronidation (~1.7 nmol APAP- Gluc/min/mg protein) and sulfation (~15.6 pmol APAP-Sulf/min/mg protein). Together, these data strongly suggest that the enhanced susceptibility of Vnn1 knockout mice to APAP hepatotoxicity is not due to differences in APAP metabolism. Immunohistochemistry of formalin-fixed liver sections following APAP treatment revealed a lack of PCNA positive hepatocytes and F4/80 positive macrophages in and around areas of centrilobular necrosis in Vnn1 knockout at 48 hours after APAP qRT-PCR from total RNA isolated from whole livers indicated that inducible nitric oxide synthase (iNos) and interleukin-4 were reduced by 2.7 fold lower in Vnn1 knockout mice at 48 hours after APAP treatment in comparison to wild-types. Myeloperoxidase and iNos exhibited a trend of decreased expression in Vnn1 knockout at 48 hours, but these differences were not statistically significant. Together, these results indicate that a lack of Vnn1 expression may alter the normal compensatory repair and immune responses following toxic APAP exposure although it is unknown to what extent these mechanisms contribute to the enhanced susceptibility of Vnn1 knockout mice to APAP hepatotoxicity.

666  Transcription Regulation by Novel Interaction of Kruppel-Like Factor 6 with Aryl Hydrocarbon Receptor at the NC-XRE.

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The Aryl Hydrocarbon Receptor (AhR) is a ubiquitous, ligand mediated basic helix-loop-helix transcription factor of Per/ARNT/Sim family that regulates adaptive and toxic responses to variety of chemical pollutants such as polycyclic aromatic hydrocarbons (PAH) and halogenated aromatic hydrocarbons, most notably 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD). Upon ligand activation, the AhR translocates into the nucleus and binds to DNA at the XRE along with aryl receptor nuclear translocator (ARNT) and facilitates transcytosis of cytochrome P450 family members. In addition, AhR-regulated gene expression is known to regulate G1 phase cell cycle progression. Recent DNA microarray experiments by our group showed that plasminogen activator inhibitor 1 (PAI-1) is a TCDD responsive gene lacking the XRE. Subsequent experimentation identified a novel non-consensus XRE (NC-XRE) located within the PAI-1 promoter that supports direct binding of AhR independent of ARNT. While the XRE and NC-XRE are distinct entities, the NC-XRE shares marked homology with the DNA binding sequence of the Kruppel-like Factor (KLF) family of transcription factors. The KLF superfamily of proteins are related to Sp1 and characterized by three zinc finger domains in their C-termini that confer binding to a GC-box linked to diverse target genes. Our results indicate that AhR interacts directly with KLF6 at the NC-XRE in a dioxin-dependent manner. Furthermore, sequential deletion studies demonstrate the importance of the AhR C-terminal and KLF6 N-terminal domains in this interaction. Since KLF6 is a known transcription factor of the G1 cyclin-dependent kinase inhibitor p21cip1, the potential exists for overlap or common roles involving both proteins in cell cycle regulation. Our further studies will focus on mapping the AhR complex at NC-XRE and understanding the role of this complex in transcription regulation. This work was supported by R01ES007800 and P30ES066676.

667  Mild Endoplasmic Reticulum Stress Preconditioning Attenuates Methylmercury-Induced Cellular Damage by Inducing Favorable Stress Responses.

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Methylmercury (MeHg) toxicity is a continuous environmental problem to human health. Failure to protect cells against MeHg-induced early oxidative stress triggers subsequent endoplasmic reticulum (ER) stress and apoptosis. Here, we demonstrate the protective effects of mild ER stress preconditioning against MeHg toxicity on a MeHg-susceptible cell line. Cells preconditioned with low concentrations (0.1-0.3 μg/ml) of an inhibitor of ER Ca2+-ATPase, thapsigargin (TPG), showed resistance to MeHg cytotoxicity through several favorable stress responses, which included phosphorylation of eukaryotic initiation factor 2 alpha (eIF2α), attenuation of activating transcription factor 4 (Atr4), and upregulation of stress-related proteins (glucose regulated protein of 78 kDa (Grp78) and metallothionein 1 (Mt1)). Atr4 accumulation was mediated by translation inhibition of its upstream open reading frame (uORF) and translation facilitation of its protein-coding ORF by the phospho-eIF2α-mediated general suppression of translation initiation, resulting in accumulated Atr4 mRNA but not protein. Integrated stress responses led to a delay of MeHg-induced oxidative stress and the activation of extracellular signal-regulated kinase pathways to promote cell survival in preconditioned cells exposed to MeHg. Finally, knockdown experiments demonstrated that Grp78 plays a crucial role in protecting preconditioned cells against MeHg cytotoxicity. These results suggested that mild ER stress preconditioning is a useful therapeutic intervention against MeHg toxicity, the underlying mechanism being the induction of integrated stress responses.

668  Differences in Regulation of Gene Expression Profiles of the Bone Marrow between C57BL/6 and C3H/Hc Mice after Benzene Treatment.

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We focused on the differences in the regulation in gene expression profiles of the bone marrow, which predict benzene-specific early responses related to plausible leukemogenic changes, between C57BL/6 (B6), a lymphoid-neoplastic-prone mouse strain, and C3H/He (C3H), a myelogenous-leukemia-prone mouse strain. Previously, we showed a latent potential of induction of myelogenous leukemia by benzene in both strains, but with different mechanistic propensities. Consequently, reciprocal differences between B6 and C3H mice in gene expression profiles related to...
669 Biphasic Influence of Arsenic, Cadmium, Mercury, and Nickel on Aryl Hydrocarbon Receptor (AHR) Signaling.

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Interactions of oxidants with the metabolic turnover of the endogenous AHR ligand 6-formylindolo[3,2-b]carbazole (FICZ) seems to play critical roles in downstream AHR-mediated signaling. Production of superoxide leading to increased proliferation was observed after exposure of immortalized human keratinocytes (HaCaT) to low levels of As3+, Cd2+, Hg2+ and Ni2+. Higher concentrations lowered the intracellular GHSH levels and led to temporal inhibition of FICZ-stimulated cytotoxicity P450A1 (CYP1A1) activity and increased adaptive antioxidant responses as measured by expression of antioxidant genes. After addition of FICZ (150 μM) together with the metals a temporal inhibition of the metabolic degradation of FICZ occurred, as determined by HPLC analyses. At these inhibitory concentrations, the metals themselves activated CYP1A1 at late time points. Transcriptional activation of CYP1A1 was estimated with a luciferase reporter assay in human hepatoma HepG2-XRE-Luc cells and the ability of the metals to activate the AHR only in the presence of the natural ligand FICZ was established by comparing AHR activation in commercial DMEM medium and in DMEM medium free from background levels of FICZ. NADPH oxidase (NOX) activity was critical for the stimulation of proliferation by the metals as determined by lack of proliferative responses in human X-CGD myeloid cells carrying a mutated gp91phox gene. In addition, influence of NOX on CYP1A1 gene expression was induced by lower metal-stimulated luciferase activity in HepG2 cells treated with the NOX inhibitor diphenyliodonium. Pretreatment with the metals led 24 h later to elevated GHSH levels and increased sensitivity of the AHR to low levels of FICZ (supershutdown). We propose that the AHR/NOX-dependent auto-regulatory feed-back mechanism, which ensures low steady-state levels of FICZ, works in close concert with the NADPH oxidase pathway.

670 Differential Gene Expression Responses by Hepatic Liver in Fischer Rats Treated with Carcinogens Phenobarbital and Diethylnitrosamine.

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Rodent liver is a primary organ for assessing compound toxicity. Non-genotoxic carcinogens such as phenobarbital (PB) and genotoxic carcinogens such as diethylnitrosamine (DEN) induce morphological and physiological changes in the liver that may vary by dose. For example, DEN induces more tumors in the right liver lobe than in the left and median lobes. To elucidate the molecular mechanisms underlying the differential cytotoxic responses, as well as the differences among the three liver lobes, groups of male Fischer rats were treated with 0 or 7.5 mg/kg/day of DEN, or 0 or 65 mg/kg/day of PB, by daily oral gavage for two and four weeks. Samples of the right lateral (RL), right median (RM), and left lateral (LL) hepatic lobes were obtained for gene expression analysis and pathological assessment. DEN induced slight apoptosis/necrosis of individual hepatocytes in all lobes, while administration of PB was associated with centrilobular hypertrophy that was much less prominent in the left lateral lobe compared with the other lobes. Gene expression profiles were assessed for three animals/group/interval using whole genome microarrays. The data showed that PB and DEN both induced robust, yet quite distinct, transcriptional changes in the animals, which could partially explain the differential morphological/physiological outcomes. In both groups of vehicle-treated rats, the transcriptional profile of the LL lobe was much different from those of the RL and RM lobes, whereas the latter two were quite similar to each other. In animals given DEN or PB, such lobe difference was much reduced, slightly more so in the rats given DEN than in those given PB. Overall, the gene expression profile may provide clues to the mechanism of PB- and DEN-induced liver tumors, and possible molecular biomarkers for predicting liver toxicity.

671 In Vitro Effects of Aldehydes Present in Tobacco Smoke on Gene Expression in Human Lung Alveolar Epithelial Cells.

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Tobacco smoke consists of thousands of harmful components. A major class of chemicals found in tobacco smoke is formed by aldehydes, in particular formaldehyde, acetaldehyde and acrolein. The present study investigates the gene expression changes in human lung alveolar epithelial cells upon exposure to formaldehyde, acrolein and acetaldehyde at sub-cytotoxic levels. We exposed A549 cells in vitro to aldehydes and non-aldehyde chemicals (nicotine, hydroquinone and 2,5-dimethylfuran) present in tobacco smoke and used microarrays to obtain a global view of the transcriptomic responses. We compared responses of aldehydes with that of the non-aldehydes. Formaldehyde gave the strongest response; a total of 66 genes was greater than 1.5-fold differentially expressed mostly involved in apoptosis and DNA damage related processes, followed by acetaldehyde (57 genes), hydroquinone (55 genes) and nicotine (8 genes). Acrolein and mixture gave effects to oxidative stress genes, no gene expression effect found on the exposure to 2,5-dimethylfuran. Overall, aldehyde responses are primarily indicative for genotoxicity and oxidative stress. These responses may relate to respiratory diseases such as cancer and COPD, respectively. The present findings could be important in providing further understanding of the role of aldehydes emitted from cigarette smoke in the onset of pulmonary diseases.

672 Regulation of the Metabolic Switch between Ductal Carcinoma In Situ (DCIS) and Invasive Breast Cancer (IBC).

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Ductal carcinoma in situ (DCIS) accounts for 15-25% of breast cancers. However, there is a gap in our understanding of the factors that regulate the progression of DCIS to invasive breast cancer (IBC). Deregulation of cell metabolism is a defining feature of tumor cells, which utilize glycolysis instead of oxidative phosphorylation. Since altered metabolism promotes invasion and metastasis, the mechanisms regulating this switch could be critical in mediating the transition from DCIS to IBC. Singleminded-2s (Sim2s) is present in early stage DCIS lesions and lost in IBC. Re-establishment of Sim2s in the MCF10DCIS.com model inhibits tumor growth, invasion, and metastasis. We hypothesize that Sim2s plays a key role in modulating the switch between DCIS and IBC. To determine the effect of Sim2s on metabolism in DCIS progression, we analyzed Sim2s over- and under-expressing MCF10DCIS.com cells for energy dependent differences in growth, autophagy, and metabolism. Live cell imaging and TEM were used to determine Sim2s’ effect on mitochondria and autophagy. Nuclear Magnetic Resonance and Seahorse analysis was employed to compare differences in metabolic signatures and measure oxidative phosphorylation and glycolytic capacity. Re-establishment of SIM2s induced oxidative phosphorylation and adaptation to metabolic stress through increased oxygen consumption, whereas loss of SIM2s promoted glycolysis by elevated glucose uptake and lactate production. These results suggest that SIM2s plays a critical role in DCIS progression and loss of SIM2s is required for the switch from oxidative phosphorylation to glycolysis.
Hypoxia Perturbs PCB-Induced Aryl Hydrocarbon Receptor Signaling and CYP1A1 Induction in Human Cells.

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The aryl hydrocarbon receptor (AhR) pathway controls cellular responses to exposure of foreign substances by activating genes that aid in xenobiotic metabolism. The ligand-activated AhR forms a heterodimer with its binding partner aryl hydrocarbon receptor nuclear translocator (ARNT) and subsequently induces the transcription of genes such as CYP1A1, a member of the cytochrome P450 family and a phase I detoxifying enzyme. Notably, ARNT proteins also dimerize with hypoxia-inducible factors (HIFs) which mediate the cellular response to low oxygen (hypoxia). During hypoxia, HIFα:ARNT heterodimers form and activate the transcription of genes that promote cell survival in low oxygen environments. Since HIFα and AhR share a common subunit, ARNT, the possibility for signaling crosstalk exists at this axis. We hypothesized that hypoxic conditions cause HIFα to sequester ARNT and thus inhibit the activation of a robust AhR transcriptional response in response to a polychlorinated biphenyl (PCB) AhR ligand. To test this hypothesis, we queried several human cell lines for their responses to PCBs. Our results indicate that CYP1A1 mRNA expression was induced by 3.3, 4.4, 5-pentachlorobiphenyl (PCB 126) in an AhR-specific manner. Exposure of the cells to hypoxia (1% oxygen for 8 hours) significantly inhibited the induction of CYP1A1 mRNA by up to 74%. EMSA and CYP1A1 promoter:luciferase reporter assays to measure DNA binding and transcriptional activities of AhR complexes further showed an inhibitory effect of hypoxia treatment on PCB-induced AhR signaling. Growth curve and MTT assays demonstrated significant growth inhibition following PCB 126 treatment. Taken together, these findings indicate that hypoxia significantly interferes with AhR-mediated responses to PCBs. Our future studies will investigate whether the observed effects also occur at the protein level and whether hypoxia can prevent the growth inhibitory effect of PCB treatment. (Supported by grant NIEHS P42 ES 013661.)

Characterizing Chronic Nicotine Exposure-Induced Behaviors and Gene Regulations in Caenorhabditis elegans.


Nicotine is one of the most abused substances. Nicotine binds nicotinic acetylcholine receptors (nACHRs), stimulates the mesolimbic dopamine system, and leads to addiction. With chronic exposures to low concentrations of nicotine are a common scenario, this study aims to characterize chronic nicotine exposure-induced behaviors and explore affected gene pathways in the model organism, Caenorhabditis elegans (C. elegans). C. elegans were treated with control, 6,17M, and 61.7 μM of nicotine for 24-hour and locomotion behaviors were characterized using a worm tracking system. Our preliminary data has determined several addictive behavioral patterns following chronic low range micromolar nicotine exposures, including stimulus, withdrawal, and tolerance. Locomotive behaviors were stimulated in control worms that were exposed to nicotine. Withdrawal behaviors were observed as increased locomotion speeds when nicotine dosages were disrupted in nicotine-dependent worms. We also detected the expressions of 39 functional genes implicated in cholinergic signaling, locomotion, egg-laying, and stress-response. Findings showed that gene expressions are most active in low but not high nicotine concentrations. The expression profiles of all the tested protein-coding genes were affected with a range from 12.5 fold down-regulation (old-1) to 138.1 fold up regulation (hid-14). Nicotine exposure also affected the expression of microRNAs, an extensive class of a small regulatory RNAs.

TNF1 Regulates the Cell Stress Response through Repression of Heat Shock Proteins A6 and A1A.


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TNF1 protein represses gene expression otherwise activated by NFkB, RAR, and PPAR. Additionally, TNF1 expression is increased in several cutaneous inflammatory diseases and wound healing. Thus, while some transcription factor targets and pathologies have been recognized, the spectrum of genes possibly affected by TNF1 is unknown. To identify additional gene expression changes due to increased TNF1 levels, we conducted a microarray analysis from keratinocytes transiently overexpressing TNF1. In addition to repression of genes known to be activated by NF-kB, RAR, and PPAR, a surprising result of decreased HSP gene family expression was observed (ex. 20x and 3.2x for HSPA6 and A1A, resp.). This reduction was confirmed by qPCR and western blot. Both HSPA6 and A1A have similar functions in cell protection and downregulation of either HSP reduced cell viability post-cell stress. Since HSPA6 basal and induced expression was decreased by TNF1, it was chosen for further studies. To guide analysis of how HSPA6 may be regulated by TNF1, we isolated and performed in silico analyses on the HSPA6 3’-kb promoter. Putative PPAR, RAR and NF-kB sites were found. PPARz and RAR ligands had no effect, but PPAR β/δ and γ ligands increased HSPA6 expression 4x. To search for TNF1-responsive region(s), we cloned 5’-HSPA6 promoter deletions into a reporter construct. Intriguingly, the most repressed region lacks PPREs, RAREs, and NF-kB sites. These results suggest that although PPAR can regulate HSPA6 and TNF1 may be repressing HSPA6 through this transcription factor, there is an additional TNF1-sensitive region independent of its previously identified PPAR, RAR and NF-kB targets. Moreover, our data suggests TNF1-mediated HSPA6 repression is independent of PPAR, RAR or NF-kB, suggesting there may be additional, as yet unrecognized, targets where interaction with TNF1 results in reduced transcriptional activity.

Erk1/2 Pathway Inhibition Attenuates BoNT/A-Induced Neurite Outgrowth in Motor Neurons.

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Botulinum neurotoxin A (BoNT/A) is known to be neurotoxic due to its paralytic effect at cholinergic synapses with extremely high potency. Independent of that, BoNT/A has been found to stimulate neurite outgrowth of motor neurons both in vivo and in vitro, which may account at least in part for the initiation of recovery from botulism. The signaling pathways regulating the BoNT/A-induced neurite outgrowth process have not been identified. In the current study, we used pharmacologic kinase inhibition to investigate the more relevant outgrowth pathways. GFP positive motor neurons were directly differentiated from HBGC3 mouse embryonic stem cells, and treated with a MEK1/2 inhibitor in the presence of 1mM BoNT/A. Following 6 and 24h post BoNT/A exposures, single motor neurons were imaged. The total neurite length and number of branches of primary and secondary neurites were measured and counted. Significant differences were determined using one-way or two-way ANOVA with post hoc tests. Preliminary results showed that 1mM of the MEK1/2 inhibitor specifically blocked BoNT/A's stimulatory effect on secondary branching at 6h compared to toxin only treated controls, suggesting an early, transient role in BoNT/A-induced signal transduction. In addition, there was no effect of MEK1/2 inhibition on the toxin's enzymatic action to cleave SNAP-25. Collectively, these data suggest that ERK1/2 pathway may play an important role in the early phases of BoNT/A-induced neurotsgenesis.

Inhibition of TGFbeta-Induced Integrin- Growth Receptor Cooperative Signaling by Curcubitamin B, Causes Breast Cancer Growth Suppression.

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Integrins have recently been shown to play an important role in promotion of cell motility and survival in various cancers through cooperative signaling with EGFRs. Integrins transmit cell survival signals through ILK-Paxillin complex and further play role in activating AKT signaling. Curcubitamin B (CuB) is a steroidal class of chemical present in the plants belonging to cucurbitaceae family. CuB has shown cytotoxic potential in several cancers by causing G2/M cell cycle arrest and inhibition of JAK/STAT3 pathway. However, the exact mechanism of CuB and its role in metastasis and cell survival is not known. Our results show that CuB significantly suppressed the growth of MDA-MB-231, SKBR3, MCF-7, MDA-MB-231 (HER2) and MCF-7 (HER2), five different breast cancer cells in a concentration and time-dependent manner. CuB also induced apoptosis and down-regulated ILK1 expression. In addition, CuB down-regulated ITGB4 and ITGα6 protein expression and suppressed the phosphorylation of AKT at Ser473, paxillin at Tyr118. Previous studies have shown that TGFβ mediates physical association of ITGB4 and HER2, through Src kinase to enhance cell survival and motility. Interestingly we observed significant downregulation of Src, EGFR and HER2 in different breast cancer cells, suggesting that CuB suppresses cooperative signaling of Integrins and HER2. We also observed that TGFβ pretreatment imparted protection from CuB's cytotoxic effects, indicating that CuB modulates TGFβ mediated Integrin signaling. Taken together our results indicate the anti-cancer effects of CuB in breast cancer cells by targeting ITGβ4/ITGα6, Src, ILK and TGFβ. Further detailed mechanistic work is in progress. [Supported in part by R01 grants CA106953 and CA210038 (to S.K.S) awarded by the National Cancer Institute].

Key: ITGB4 – Integrin β4, ITGα6 – Integrin 66, TGFβ – Transforming growth factor β, ILK – Integrin linked kinase, EGFR – Epidermal growth factor receptor, CuB – Curcubitamin B
Drug-induced liver injury (DILI) is an important clinical problem and predicting human DILI for novel candidate drugs is difficult. Current models indicate that in many cases DILI is linked to reactive metabolite formation and involves activation of innate immune systems. Here we systematically evaluated the combined application of high content live cell imaging-based analysis of 1) NRF2 activation as a measure for reactive metabolite stress; 2) perturbations of the normal NF-κB activation mediated by TNG6; 3) synergistic induction of apoptosis by DILI compounds and TNFα. Fifteen drugs associated with DILI were evaluated. Most DILI compounds induced NRF2 stabilization-dependent activation of the downstream target SRXN1 (11 out of 15). Various DILI compounds affected the TNFα expression in association with protection against the drug/TNFα-induced apoptosis independent of drug-induced oxidative stress. Interestingly, CBZ decreased Chk1 expression, whilst simultaneously inducing Ser-345 Chk1 phosphorylation. RT-PCR data revealed that MG-132 down-regulates Chk1 gene expression. Finally, following S-phase cell synchronization with aphidicolin, TGHQ strikingly attenuated the S-phase arrest, which may drive HL-60 cells into apoptosis in HL-60 cells. We now report that TGHQ induced severe DNA damage, as evidenced by DNA ladder formation and histone H2AX phosphorylation, and the subsequent engagement of checkpoint response pathways. Thus, TGHQ induced the activation of both ATR and ATM, which initiate the mammalian DNA damage response via the phosphorylation of the effecter proteins Chk1 and Chk2. Perturbation of Chk1 signaling abrogates the TGHQ-induced S-phase arrest, indicating an important role for Chk1 in the S-phase checkpoint following TGHQ-induced DNA damage. Moreover, Cdc25A, which is required for the intra-S phase checkpoint, was degraded in HL-60 cells treated with TGHQ. TGHQ also substantially decreased Chk1 levels, primarily by decreasing the Chk1 gene transcription. Consistent with these findings, MG-132, a proteosome inhibitor, failed to restore Chk1 protein levels. Interestingly, MG-132 itself (> 0.1 μM) decreased Chk1 expression, whilst simultaneously inducing Ser-345 Chk1 phosphorylation. RT-PCR data revealed that MG-132 down-regulates Chk1 gene expression. Finally, following S-phase cell synchronization with aphidicolin, TGHQ strikingly attenuated the S-phase arrest, which may drive HL-60 cells into premature mitosis in the presence of un-repaired DNA damage. In summary, TGHQ-induced DNA damage triggers the ATM/ATR-Chk1/Chk2-Cdc25 pathway and induces Chk1-dependent S-phase arrest in the p53-deficient HL-60 cells. The results are consistent with the view that targeting of Chk1 is an effective chemotherapeutic strategy. However, the data also reveal that in addition to the established mechanism of Chk1 protein degradation engaged by the majority of anti-cancer agents, a novel mechanism to modulate Chk1 protein expression can be engaged at the transcriptional level.
factor tool predicted the activation of SREBF1 and SREBF2 comprising the induc-
ction of cholesterol uptake (Ldlr), cholesterol biosynthesis (Hmgcr, Hmgcs1, Lss,
Sgce, Cyp51) and cholesterol transporter (Star) gene expression. Additionally, the
predicted activation of CREBF1, CREBF2, CEBPA and CEBPB suggest increased
CAMP-mediated gene expression such as the induction of Vegfa, Gadd45a, and
Csf1 involved in cell morphology and growth. Collectively, these results suggest
the late effects of ATR on BLTK1 murine Leydig cell steroidogenesis are mediated
by SREBF- and CREB-mediated changes in gene expression.

683 Hepatic Gene Expression Analysis of 2-Aminoanthracene-
Exposed Fisher-344 Rats Reveal Patterns Inductive of Liver
Carcinoma and Type 2 Diabetes.

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The goal of the present study was to examine hepatic differential gene expression
patterns in Fisher-344 rats in response to dietary 2-aminoanthracene (2AA) inges-
tion for 14 and 28 days. Twenty four post-weaning 3-4 week old F-344 male rats
were exposed to 0 mgkg-1-diet (control), 50 mgkg-1-diet (low dose), 75 mgkg-1-
diet (medium dose) and 100 mgkg-1-diet (high dose) 2AA for 14 and 28 days. This
was followed by analysis of the liver for global gene expression changes. In both
time points, the numbers of genes affected seem to correlate with the dose of 2AA.
Sixteen mRNAs were differentially expressed in all treatment groups for the short-
term exposure group. Similarly, 51 genes were commonly expressed in all 28-day
expression group. Almost all the genes seem to have higher expression relative to the
controls. In contrast, cytochrome P450 family 4, subfamily a, polypeptide 8
(Cyp4a8), and monocyt to macrophage differentiation-associated (Mmd2) were
down-regulated relative to controls. Differentially expressed mRNAs were further
analyzed for associations via DAVID. GO categories show the effect of 2AA to be
linked with genes responsible for carbohydrate utilization and transport, lipid
metabolic processes, stress responses such as inflammation and apoptosis processes,
immune system response, DNA damage response, cancer processes and circadian
rhythm. The data from the current study identified altered hepatic gene expression
profiles that may be associated with carcinoma, autoimmune response, and/or type 2
diabetes. Possible biomarkers due to 2AA toxicity in the liver for future study in-
clude Abcb1a, Nhej1, Adam3, Cdkln1, Mgmt, and Nenc.

684 Utilizing RNA Sequencing and the DNA-Damage Response
 to Model Chemical Toxicity: How PKC-Activating Tumor Promoters Create a Path of Their Own.

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The DNA damage response (DDR) is an essential cellular pathway for maintaining
genomic integrity and preventing neoplastic transformation. Although tumor pro-
moting compounds do not directly cause fixed mutations, they have been previ-
ously shown to alter many different cellular processes that also impact normal
DNA damage signaling. Therefore, the purpose of this study was to profile PKC activating
tumor promoting chemicals by their respective alterations to normal DDR signal-
ing induced by short-wavelength ultraviolet light (UVC). Human lymphoblastoid
TK6 cells were exposed to 1 nm 12-O-tetradecanoyl-phorbol-13-acetate (TPA) for
72 hours which caused a reduction in cell growth and changes in cellular morphol-
ogy. Following UVC exposure (10 J/m2), TPA treated cells displayed a synergistic
increase in apoptosis and a significant delay in γH2AX clearance. Other PKC activ-
ing tumor promoters including other phorbol esters, mezerein, sapintoxin D, (−)
-indolactam V, and resiniferonol 9,13-ortho-phenylacetate caused similar
changes in cell morphology and a synergistic increase in apoptosis following UVC
exposure. RNA sequencing was used to capture the global transcriptomic profile 8
hours after UVC-induced DNA damage following TPA pretreatment. Principle
component analysis of UVC treated cells revealed a relationship between gene ex-
pression patterns and the concentration of TPA (0.2, 0.5 and 1 nM). These find-
ings indicate a deviation from the normal DDR response. TPA pretreatment at 1
nM caused greater than 300 of the UVC-regulated genes to be downregulated and
greater than 80 UVC-regulated genes to be upregulated compared to the normal
UVC expression profile. These results show that the tumor promoter TPA has a
unique gene expression profile following DNA damage that can be potentially used
to profile this class of chemical.

685 Reactive Nitrogen Species Regulate Autophagy through
ATM-AMPK-TSC2-Mediated Suppression of mTORC1.

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Reactive intermediates such as reactive oxygen species (ROS) and reactive nitrogen
species (RNS) play essential roles in the cell as signaling molecules, but in excess,
constitute a major source of cellular damage in both physiological (i.e. mitochondr-
ial respiration) and pathophysiological (i.e. inflammation) settings. Recently, ROS
were shown to induce autophagy via a novel ATM signaling pathway in the cyto-
plasm. Activation of cytoplasmic ATM by ROS engages LKB1 and AMPK kinases
to activate the TSC2 tumor suppressor, which represses mTORC1 (a negative reg-
ulator of autophagy) to induce autophagy. However, whether RNS engage this or
similar signaling pathway(s) to regulate mTORC1 and induce autophagy is unclear.
We found under conditions of steady-state nitric oxide (NO) exposure, rapid aci-
tivation of ATM and downstream signaling by this stress kinase to LKB1-AMPK and
the TSC2 tumor suppressor. As a result, mTORC1 was repressed as evidenced by
decreased phosphorylation of the direct mTORC1 targets S6K, 4E-BP1, and ULK-
1, the target for mTORC1 repression of autophagy. Induction of autophagy con-
cordant with decreased ULK1-1 phosphorylation by mTORC1 at S757 and in-
creased phosphorylation at S317 by AMPK was demonstrated by increased ratio
of LC3-II/LC3-I, formation of GFP-LC3 puncta, increase in acidic vesicles and de-
crease p62 levels, indicative of increased autophagic flux. While autophagy has
been shown to have pro-death and pro-survival functions, induction of autophagy
by NO caused loss of cell viability, suggesting it functions primarily as a cytotoxic
response to excess nitrosative stress. These data indicate that nitrosative stress, like
oxidative stress, regulates autophagy. As cancer cells are particularly sensitive to ni-
troxative stress, these data open new therapeutic avenues capitalizing on the ability
of NO and RNS to induce autophagic cell death.

686 CYP2S1 Expression Influences Cytotoxicity of the
Anticancer Prodrug, AQ4N, in Human Bronchial Epithelial
Cells.

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Cytochrome P450 2S1 (CYP2S1) is one of the most upregulated P450s identified
in human cancers and correlates with poor prognosis. The underlying cause of ele-
vated CYP2S1 expression and the consequence of elevated expression is not
known. The only recognized substrate for CYP2S1 is AQ4N (Banantranone). AQ4N is
a bioreductive prodrug under clinical investigations for hypoxic tumors. AQ4N is
metabolically reduced in hypoxic environments to the topoisomerase II inhibitor,
AQ4. CYP2S1 is recognized as one of the most efficient enzymes to catalyze this re-
duction. The objective of this study was to determine: i) whether human CYP2S1,
like the mouse CYP2s1, is elevated in response to hypoxia; and ii) whether changes
in CYP2S1 expression influence CYP2S1-mediated metabolism of AQ4N. To
determine whether CYP2S1 was induced in response to hypoxia we evaluated mRNA
expression in the presence of 1% oxygen as well as the hypoxia mimetics cobalt
chloride and α-phenanthroline. CYP2S1 mRNA was significantly increased by ap-
proximately 8-, 3-, and 50-fold in response to 1% oxygen, cobalt chloride, and α-
phenanthroline. These results suggest that human CYP2S1 mRNA is elevated in re-
sponse to hypoxia. To determine whether changes in CYP2S1 expression influence
AQ4N metabolism we examined the effects of AQ4N and AQ4 on cell viability in
bronchial epithelial cells (BEAS-2B) cells differentially expressing CYP2S1.
Cytotoxicity was estimated using alamar blue reduction in cells expressing high
(CYP2S1-FLAG), medium (control plasmids: pcDNA3.1 and scrambled shRNA),
and low CYP2S1 expression (CYP2S1 shRNA). Preliminary results demonstrate that
elevated CYP2S1 expression does enhance cytotoxicity under hypoxic condi-
tions (0.2%-1% O2). Interestingly, our results also suggest a protective role for
CYP2S1 in normoxic conditions (21% O2). These data suggest that elevated
CYP2S1 expression may have a dual role in protecting normal cells while promot-
ing AQ4N-mediated cytotoxicity in hypoxic cells.
687 Transcriptome Analysis Reveals Novel Pathway Regulation in Response to Differential Expression of CYP2S1 in Human Lung Cells.

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Cytochrome P450 2S1 (CYP2S1) is considered an orphan P450 with an unknown biological function. Data from our laboratory and others suggest that CYP2S1 may have an important physiological role in modulating the synthesis and metabolism of bioactive lipids, including prostaglandins and retinoids, respectively. CYP2S1 expression is elevated in epithelial-derived cancers. Whether increased CYP2S1 expression in proliferative disease is protective, detrimental, or fails to impact disease progression remains to be determined. To elucidate its role, we need to understand the physiological significance of CYP2S1. We reasoned that transcriptome analysis of human bronchial epithelial cells (BEAS-2B) differentially expressing CYP2S1 would reveal metabolic shifts in key regulatory pathways linked to CYP2S1-mediated metabolism. To test this idea, we established four stable BEAS-2B cell lines expressing high (CYP2S1-Flag), medium (pCDNA3.1 and scrambled shRNA), and low (CYP2S1 shRNA) CYP2S1 expression. Expression levels confirmed using qRT-PCR and western analysis. Alterations in the transcriptome were determined using RNA-sequencing (RNA-seq) analysis. RNA was isolated from three biological replicates from each of the four experimental conditions (CYP2S1-Flag vs pCDNA3.1 and CYP2S1 shRNA vs SCRAM shRNA). Eight to fourteen million sequencing reads were generated from each of the 12 samples. Four distinct population clusters were identified using principal component analysis, representing each of the transduced cell lines. Approximately 1000 genes were differentially expressed in response to CYP2S1. Among these were key regulatory enzymes involved in prostaglandin synthesis. RNA-seq also identified novel changes in the mTOR pathway, which regulates cell size. Significant differences in cell size was in lung cells differentially expressing CYP2S1, revealing a potentially novel role for CYP2S1 in cell size regulation.

688 In-Depth Examination of How Exposure to Long-Term Low-Dose Chromium Alters the Transcriptional Response to BaP.

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Complex mixtures and their toxicity are a significant and growing problem that must be addressed by applied toxicology. Techniques for the assessment of similar compounds in a mixture based on relative toxicity are inadequate for the assessment of complex mixtures with different types of components. In complex mixtures, multiple exposures generate specific changes in gene expression that cannot be attributed to any single mechanism. Data on the responses to these kinds of mixtures are rare and often contradictory. Our previous work demonstrated that epigenetic regulation of gene transcription plays an important role in chromium's toxic response, and that Cr is capable of altering the transcriptional regulation of known BaP responsive genes, like Cyp1a1. Here we use RNA-seq after long-term low-concentration exposure to chromium to examine the extent to which chromium exposure can alter BaP induced transcriptional responses throughout the transcriptome. We find that mouse hepatoma cells exposed to long-term low concentrations of sodium chromate, in the range of 0.1 – 0.5 μM, accumulate double-strand DNA breaks, and that the percentage of cells undergoing apoptosis increases concurrently with either increased concentration or longer period of exposure to Cr. Using RNA-seq we determined that the transcription of genes critical to several important processes are regulated differently by BaP if the cells were previously chronically exposed to a low concentration of Cr. In cells grown normally genes related to critical processes are regulated differently by BaP if the cells were previously chronically exposed to a low concentration of Cr. Through these studies we hope to develop a better understanding of how exposure to chromium alters the transcriptional response to BaP.

689 Expression of the Nuclear Receptor Corepressor Tnip1 in Mouse Tissues.


Tnip1 regulates multiple signaling pathways (NF-κB, ERK2, PPAR and RAR) involved in either executing or responding to toxic insults. Altered Tnip1 expression levels are associated with certain human inflammatory diseases. These cell signaling and pathology studies revealed some molecular targets and potential organ end-points. Additionally, body-wide data on expression of Tnip1 is incomplete. To develop a better understanding of Tnip1’s expression pattern and functional role, we developed a Tnip1 gene trap knock out (KO) mouse to identify possible new signaling and organ targets.

The Tnip1 gene trap KO mouse was created via a β-Geo cassette (β-galactosidase and neomycin resistance genes) insertion into the Tnip1 first intron. Expression from the cassette allowed for evaluation of Tnip1 promoter activity in different tissues qualitatively via X-gal staining, immunofluorescent (IF) staining for β-Galactosidase (β-Gal), and quantitatively via Q-PCR. With WT tissues as a control, all heterozygote (Tnip1 +/-) tissues turned blue with X-gal staining. Results of IF β-Gal staining in Tnip1 +/- mouse tissues were similar to that of X-gal whole organ staining and paralleled the Tnip1 +/-body staining. For most organs, both β-Gal and Tnip1 were homogeneously expressed across the tissue. However, for kidney, both β-Gal and Tnip1 glomerular staining was significantly reduced compared to convoluted tubules suggesting distinct expression levels and fidelity of the β-Gal marker for the endogenous Tnip1 expression. Q-PCR results revealed that Cr specifically altered transcriptional activity as determined by relative levels of β-Gal mRNA, varied among different tissues and mirrored the Tnip1 mRNA levels. These results suggest Tnip1 is expressed at varying levels in different mouse tissues. Recent studies report a high rate of lethality in Tnip1 KO mouse embryos due to fetal liver apoptosis (Oshima et al, 2009, Zhou et al, 2011). By contrast, we demonstrate there are tissues other than liver expressing Tnip1 and that biological function of these organs may be impacted by changes in levels of Tnip1 protein.

690 Evaluating the Role of Kinase Activation Involved in Chemical-Dependent p53 Response to DNA Damage.

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Post-translational modifications (PTMs) of p53, individually and in combination, help determine the function of p53 in DNA damage response, e.g. cell cycle arrest, apoptosis and DNA repair. The dose-response for p53 activation, cell cycle, apoptosis and mutation (micronuclei) were studied for etoposide (ETP), methylmethane sulfonate (MMS) and quercetin (QUE). Cellular response was chemical specific, despite similar induction of DNA damage (double strand breaks) and p53. Here, we investigated activation of mitogen-activated protein kinases (MAPKs) and phosphoinositide 3-kinases (PI3Ks) in response to MMS, ETP and QUE across doses and time (1 – 24 hrs) in human fibrosarcoma cells (HT1080). The MAPKs were studied by measuring p-p38, p-ERK1/2, p-JNK1/2, and p-MEK1 using immunoblot and Luminesin assays, and the PI3Ks: p-ATM, p-ATR, p-DNA-PK and p-Chk2 by immunoblot and flow cytometry. MAPKs and ATM were activated by all three chemicals. Although p-ATM, p-p38, p-ERK and p-JNK increased at 1, 3, and 24 hrs, they showed different patterns of activation that may lead to observed differences in cell fate. While ETP and MMS showed maximal p-ERK1/2 and p-ATM at 24 hr, QUE displayed a much stronger p-ERK and p-ATM response at 1 hr, which may indicate more efficient repair, and is in agreement with less micronuclei observed with QUE vs. ETP or MMS, p-p38 - associated with p-p53 (ser46) leading to apoptosis - was induced quickly with ETP (1 hr), but more slowly with QUE, an observation consistent with observed apoptosis by ETP but not QUE in HT1080 cells. ETP showed a much stronger induction of p-ATM than either MMS or QUE, and only QUE inhibited p-ATR. Our results indicate that, types and timings of the activated kinases are chemical specific and likely to determine the downstream actions of p53. Currently, we are evaluating treatment dependent PTMs of p53 by proteomics, as well as kinase specific inhibitors to further confirm the relationship between kinase modification of p53 and cell fate.

691 Global Transcriptional Analysis of Zebrafish Embryos following Acute Exposure to Cadmium.

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Although cadmium (Cd) exerts its toxic effects through the generation of oxidative stress, many of the cellular mechanisms involved in Cd toxicity are not fully established. Tnip1 was identified as a zebrafish embryo model, we undertook a transproteomic-based approach to investigate the role of Cd as a developmental toxicant and assist in the identification of novel mechanisms of cellular dysfunction. Zebrafish embryos (24, 48, 72 and 96 hpf) were continuously exposed to Cd (25, 50, 100, 150 μM) for 24 hours prior to mortality assessment. No significant mortality was observed prior to 72 hr. However, at 96 and 120 hpf the 50 μM Cd resulted in 20% and 40% mortality, respectively, while the 100 and 150 μM Cd doses resulted in an average of 75% and 95% mortality, respectively. To investigate global changes in gene expression, zebrafish embryos (72, 96 and 120 hpf) were exposed...
to 50 μM Cd for 4 or 8 hours. Global gene expression profiling was performed using the Agilent 4×4K Zebrafish Oligo Microarray. Overall, 778 probes were significantly upregulated, while 679 probes were downregulated in one or more treatments. Principle component analysis was employed to find trends among the various treatments. The first principle component separated out the 96 and 120 hpf Cd-treated embryos, consistent with the mortality curves suggesting that zebrafish embryos are most sensitive to Cd toxicity around 96 hpf. The majority of genes altered by Cd exposure resulted in the upregulation of a large subset of genes responsive to oxidative stress, genes involved in glutathione synthesis and heme/iron homeostasis, mitochondrial uncoupling proteins, and various solute carriers with roles in zinc transport/homeostasis or mitochondrial oxidative phosphorylation. The majority of genes downregulated by Cd were involved in cell cycle control and DNA replication. A thorough analysis of genes differentially regulated by Cd during zebrafish development will be presented. [R00ES017044]

692 Role of Chromatin Structural Changes in Regulating Human CYP3A Ontogeny. N. L. Giebel1, J. D. Shadle1, K. Dorko2, S. C. Strom1, P. M. Simpson1, K. Yan1 and R. N. Hines1, 1Pediatrics, Medical College of Wisconsin, Milwaukee, WI; 2Pharmacology, Toxicology and Therapeutics, University of Kansas Medical Center, Kansas City, KS; 1Laboratory Medicine, Karolinska University Hospital, Stockholm, Sweden.

Variability in drug metabolizing enzyme (DME) developmental trajectories contributes to differential susceptibility to chemical toxicity and adverse drug reactions, particularly in the first years of life. Factors linked to variability are largely unknown and molecular mechanisms regulating ontogeny are likely involved. To evaluate chromatin structure dynamics as a contributing mechanism, age-dependent changes in histone marks were evaluated within known CYP3A4, CYP3A7, FMO1, and FMO3 regulatory domains. Chromatin immunoprecipitation with fetal and adult primary human hepatocyte chromatin pools followed by qPCR was used to determine relative histone mark occupancy. Histone mark occupancy is consistent with many regions of bivalent chromatin (i.e., exhibiting histone marks associated with active and repressive transcription) in adult and fetal hepatocytes with the exception of the CYP3A7 XREM and C/EBPβ domains in adult hepatocytes. For CYP3A4 the bivalent histone mark domains are consistent with a low level of fetal liver expression. Differential histone mark occupancy indicates adult histone mark and transcription factor occupancy have a developmental trajectory towards an active transcriptional state. In contrast, the bivalent CYP3A7 regions in fetal hepatocytes do not correspond with high expression and surprisingly, adult hepatocytes had less histone mark occupancy, particularly the repressive mark H3K27me3. These findings for CYP3A7 suggest mechanisms other than chromatin structural changes involved in regulating ontogeny. Chromatin structural change is an important mechanism controlling DME ontogeny, particularly for those whose specific activity increases substantially after birth. Supported in part by NIH grant GM081544.

693 6-Formylindolo(3,2-b)carbazole (FICZ) Positively Regulates the Signalsome Responsible for Retinoic Acid (RA)-Induced Differentiation of Leukemic Blast Model HI-60. R. Bunaciu and A. Yen. Cornell University, Ithaca, NY.

Tryptophan is an essential amino acid that absorbs strongly in UV (extinction coefficient: 5,050 M-1cm-1 for 280 nm). Both in cell free conditions (such as cell culture media exposed to light) and in the cells (such as skin keratinocytes exposed to sunlight), tryptophan yields a multitude of photoproducts. One such photoproduct is 6-Formylindolo(3,2-b)carbazole (FICZ). FICZ has a high affinity for AhR (Kd<7x10-11 M). It is proposed to be an endogenous AhR agonist and also a cytochrome P450 family 1 substrate. FICZ and a multitude of its metabolites have been shown to be present in human fluids. Currently all the complete parental nutrition formulations contain tryptophan. The bags for total parenteral nutrition are made of clear plastic. Since many cancer patients receive both parenteral nutrition and chemotherapy (generally enhancing AhR expression and activity), it is of significant interest to assess potential effects of FICZ on cell differentiation and in particular on leukaemic induced leukemic cell differentiation. HL-60 leukaemic cells were untreated or treated with RA alone or RA in combination with FICZ for 48 h. To assess the involvement of AhR, other cells were treated either with α-tapthalnone and RA or β-naphthalonone and RA. Expression of cell surface differentiation markers, cell cycle distribution and respiratory burst response were quantified by flow cytometry. Western blots were performed to assess the proteins known to be part of the differentiation driving the induced differentiation. FICZ augments the RA-induced differentiation as evidenced by CD38 and CD11b receptors, inducible respiratory burst, G0 cell cycle arrest and growth curves. Signaling molecules previously shown to be upregulated by RA and to drive RA induced differentiation were enhanced by FICZ: MAPK signaling axis, Src family kinases, Cbl, VAV, SLP-76, P13K, and IRF-1. Moreover, AhR is associated with the c-CBL adaptor in the signalsome, as shown by IF. In conclusion, FICZ modulates RA induced differentiation.

694 Nrf2 Gene Regulation during Oxidative Stress in Embryonic Development. A. R. Timme-Laragy1,2, S. I. Karchner1, R. C. Harber1, A. G. MacArthur2 and M. E. Hahn1, 1Pediatrics, Medical College of Wisconsin, Milwaukee, WI; 2Bioengineering, University of Wisconsin, Madison, WI; 3Public Health, University of Massachusetts, Amherst, MA; 4AAMC Consulting, Hamilton, ON, Canada.

Nrf2 is a transcription factor that regulates antioxidant defenses in response to oxidative stress. Embryonic development is highly susceptible to disruption by exposure to chemicals, including those that alter redox balance. The role of Nrf2 in the oxidative stress response (OSR) during embryonic development remains unclear. Our previous work identified a novel Nrf2 paralog, nrf2b, in zebrafish (Danio rerio). This study builds upon that work to elucidate the roles of nrf2a and nrf2b in regulating the OSR during vertebrate embryonic development. Zebrafish embryos were micro-injected with antisense morpholino oligonucleotides (MO) to knock down translation of either nrf2a or nrf2b, or a standard morpholino control (co-MO). At 48 hours post fertilization, embryos were exposed to the pro-oxidant and Nrf2 activator, tert-butylhydroquinone (tBHQ) or vehicle (DMSO) for 4 hours, and preserved for RNA isolation. Microarrays were conducted using Agilent’s V3 4x4k array, and selected genes validated by QPCR. In response to tBHQ, 71 probes were up-regulated in the co-MO group, including gap1, ggc, ferritin, peroxiredoxin1, hsp70, sod1, and other genes typically found as part of the OSR. Interestingly, we found that an important and often overlooked part of the OSR is the down-regulation of genes, including cathespin, various complement components, and apolipoprotein E. Knockdown of Nrf2a or Nrf2b blocked some but not all of the tBHQ-induced changes in OSR gene expression and the effects of Nrf2a-MO and Nrf2b-MO were distinct. The results show that Nrf2 paralogs primarily regulate distinct gene sets, with some overlapping targets, in response to oxidative stress in embryos. This study also highlights the importance of gene down-regulation as a component of the OSR during embryonic development. [F32ES017585, R01ES015636].

695 On Breast Cancer Treatment, Synergistic Effects of Akt1 shRNA and Paclitaxel-Incorporated Conjugated Linoleic Acid-Coupled Poloxamer Thermosensitive Hydrogel. S. Hong1, C. Cho2 and M. Cho1, 1Laboratory of Toxicology, Seoul National University, Seoul, Republic of Korea; 2Agricultural Biotechnology, Seoul National University, Seoul, Republic of Korea.

The phosphonositoise 3-kinase(PI3K)/Akt1 signaling pathway has emerged as a target for treatment of cancer therapy. In this study, we develop a strategy to enhance Akt-targeted therapy. We thought that combination of Akt1-targeted therapy with conventional chemotherapy using paclitaxel-incorporated conjugated linoleic acid-coupled poloxamer (CLA-CP) thermosensitive hydrogel may have synergistic effects in cancer therapeutics compared with chemotherapy system. FDA-MA-231 cell, the human breast cancer cell line, was inoculated into 6-week-old female BAL/C nu/nu athymic mice. After 2 weeks of inoculation, shAkt1 and CLA-CP treatments were subcutaneously injected into the tumor. We found that the combination of shAkt1 with paclitaxel showed synergistic anti-cancer effects, thus, inhibiting the growth of human breast cancer cells, and breast cancer xenografts in mice as well. The combination therapy showed enhanced anti-cancer effects through inhibiting Akt1 signaling, inducing apoptosis. In addition, it suppressed angiogenesis and proliferation, also. It suggests that the presented strategy of combination of shAkt1 with paclitaxel may have a potential for the treatment of breast cancer. Acknowledgements This work was partly supported by the National Research Foundation (NRF-2011-000380), Ministry of Education, Science and Technology (MEST) in Korea.


Chronic Obstructive Pulmonary Disease (COPD) is a progressive disease of the respiratory system which is characterized by destruction of the epithelial cells of lungs. Current therapies alleviate symptoms of COPD but do not treat the under-
lying causes. It has been recognized in the literature that the T-box transcriptional factor TBX5 is suppressed in patients with COPD. The purpose of the study was to identify regulatory target genes of TBX5 and their significance in future treatment of COPD. A compendium of lung epithelium microarray datasets was created using GSE 19027 and GSE 994 from the Gene Expression Omnibus database. Probe IDs for TBX5 and genes that have known interactions with the TBX5 transcription factor were used to generate an initial network. Bayesian Network Inference with Java Object was then used to generate a new regulatory network, given the gene expression data. Cytoscape was used to visualize the differences between the initial and predicted networks. In addition, analysis of GSE 1650, which compared lung tissue from smokers with severe emphysema and smokers with mild or no emphysema, was done. The resulting network indicates that STAT3 regulates TBX5, NKK2-5, GATA4, ID2 and TAZ. The regulatory relationships between STAT3 and TBX5, NKK2-5, GATA4, ID2 have been validated in the published literature. However, a relationship between STAT3 and TAZ has not been established. It is known that TAZ regulates TBX5. This presents with a possible feed-forward relationship among TBX5, STAT3 and TAZ. STAT3 is known to be involved in cell growth, apoptosis, and regulation of anti-inflammatory response by controlling genes involved in the inflammatory response. This result suggests a potential inhibitory relationship between STAT3 and TBX5 since STAT3 was up-regulated in smokers with severe emphysema and TBX5 was suppressed in patients with COPD. The result suggests that future therapies targeting STAT3 or TBX5 can regulate inflammatory response involved in COPD. Further research is warranted to investigate how TBX5, STAT3 and TAZ play a role in the inflammatory response of COPD.

697 Suppressing Effect of HES1 (Drosophila Hair and Enhancer of Split-1) on Human Hepatic Multidrug Resistance-Associated Protein 4 (MRP4) Gene Expression and Inhibition of HES1 by Oxidative Stress.

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Previous in silico studies in our laboratory identified a total of 107 canonical HES1-binding N-boxes (CANNAG) in 10kb DNA fragment upstream to the translation start site of the multidrug resistance-associated protein 4 (MRP4/ABCC4). In this study, we investigated the role of HES1 in regulating MRP4 gene expression. We have identified a 16-bp DNA fragment in the MRP4 promoter-promoter proximal region containing a non-canonical N-box (GGCGTG) for HES1 that accounts for a profound suppression of MRP4 promoter transcription activity in HepG2 cells. A point mutation in this cis-element (GGGTCC) resulted in the loss of this suppressive effect of HES1. Furthermore, suppression of MRP4 reporter genes containing 230 bp, 5 kb or 7 kb of 5' flanking regulatory sequences was observed by over-expression of HES1 but not by a dominant negative mutant form of HES1 which is unable to bind to DNA. From these findings we hypothesized that the MRP4 induction we have previously reported under conditions of oxidative stress might be due in part to down-regulation of HES-1 expression. To our surprise, HES1 mRNA levels are significantly increased in the human liver cell line HC04 treated with the pro-oxidant chemicals tert-butyldihydroperoxide or tert-butyldihydroquinone (10mM). We also detected HES1 protein accumulation in nuclei of HC04 treated with these chemicals in a dose- and time-dependent manner. Similarly, treatment of mice with toxic doses of APAP increases liver Hes1 mRNA levels. These increases in expression and function of the repressive factor HES1 by oxidative stress seem inconsistent with the induction of MRP4 that also occurs under oxidative stress conditions. Studies are currently underway to investigate whether HES1 directly regulates MRP4 gene expression in vivo by oxidative stress and to determine the interplay of HES1 with other activating and repressive transcription factors known to regulate MRP4. Supported by NIH Grant DK06957.

698 Exposure to Metals Mixtures: Genomic Alterations of Infectious Disease Response Pathways in Children Exposed to Environmental Metals.

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Exposure to toxic metals can have harmful health effects, particularly in children. Although studies have investigated the individual effects toxic metals have on gene expression and health outcomes, there are no studies assessing the effect of metal mixtures on gene expression profiles. Here, we assessed the mixture effect of six toxic metals (arsenic, beryllium, cadmium, chromium, mercury, and lead) on gene expression in children in Detroit, Michigan. Indicators of Childhood Asthma (MICA) cross sectional study, we assessed metal exposure in 131 children in Detroit using fingernail metals levels. A metals mixture score was calculated and compared to gene expression profiles across the population adjusting for age and race. There were 145 unique genes that were significantly differentially expressed between children exposed to low and high levels of the metals mixture. Of the genes differentially expressed, 107 (74%) had increased expression while 38 (26%) had decreased expression. The main biological function associated with multiple metals was infectious disease. Within that group, genes were associated with infection of respiratory tract (P < 10^-6) severe acute respiratory syndrome (P < 10^-5). Taken together, the data demonstrate that exposure to metals mixtures may activate gene networks related to infectious disease response. This abstract does not necessarily reflect the views or policies of the EPA.

699 Transcriptomics Analysis of Lung Tissues of Brown Norway Rates Exposed to the Sensitizers Trimellitic Anhydride (TMA), Oxazolone (OXA) or Dinitrochlorobenzene (DNCB).

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TMA is a respiratory allergen, OXA is a potent contact allergen in man, but elevates serum IgE levels and Thelper-2 cytokines in test animals, suggesting that it has also respiratory allergic potential. DNCB is a contact allergen. Brown Norway rats (BN) were sensitised by two dermal applications of TMA, OXA or DNCB. The animals were exposed by inhalation on day 21. TMA and OXA both induced elevation of serum IgE, breathing pattern changes and the same type of allergic laryngal inflammation. However, microarray analysis of the lung (sampled 24 hrs after provocation) indicated that OXA may act through different mechanisms than TMA, despite a certain overlap in activated genes. TMA upregulated strongly Th2-associated genes and genes associated with lung remodeling, whereas OXA activated predominantly Th1-associated genes. The variability in balance between Th1- and Th2-associated genes may reflect different subtypes of respiratory allergies. DNCB induced a very mild inflammation in the larynx, which resembled DNCB-induced inflammation in the skin; only a few Th1-associated genes were differentially expressed in the lung. The transcriptomics analysis supports the idea that classification of allergens based on single genes is unreliable. Omics data contribute to our understanding of respiratory allergy and may help in the design of a predictive toxicity test for sensitization.

700 In Vivo Expression of P16INK4a in Response to Toxicological Exposures.

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In mammals, expression of P16INK4a is highly regulated. Excess expression can lead to cellular senescence and aging, while impaired activation is associated with cancer. The precise mechanism of p16INK4a regulation in vivo is poorly understood. In vitro systems have limited utility since proliferation in culture induces p16INK4a. Both extrinsic (chemotherapy and ionizing radiation) and intrinsic (telomere shortening and improper DNA damage repair) stimuli can induce p16INK4a, but the kinetics of and cellular responses to these genomic insults have not been examined in vivo. To address this question, we developed a murine strain with firefly luciferase ‘knocked-in’ to the endogenous p16INK4a locus and under control of the p16INK4a promoter (p16-LUC). We exposed p16-LUC mice to 50ppm arsenic, 42% fat diet, 350 J/m2 UVB light, or cigarette smoke for a minimum of 6 months. Every other month, p16-LUC mice were imaged to measure luciferase induction. At 30 weeks of exposure, mice exposed to cigarette smoke displayed 2 times higher levels of whole body luciferase activity than no-smoke controls. Additionally, mice exposed to UVB exhibited 1.5 times higher levels of luciferase activity by 6 weeks of exposure that reached 8 times over control mice by 24 weeks. In arsenic exposed mice, there was only a slight induction of p16INK4a by 24 weeks (not statistically significant). We observed no differences in p16INK4a expression in mice on high fat diet (42% fat) compared to normal fat diet (4%) after 78 weeks. Currently, we are correlating luciferase expression in different organs with tissue damage and mRNA expression of p16INK4a. The data generated from these experiments will demonstrate how environmental exposures are associated with expression of p16INK4a, a mediator of tumor suppression and aging. (Supported by T32 ES07126, HHMI Translational Medicine Program, and U01 CA141576)
Long Interspersed Nuclear Element-1 (LINE-1 or L1) is an autonomous, mobile element within the human genome that transposes via a ‘copy and paste’ mechanism and relies upon L1-encoded endonuclease and reverse transcriptase (RT) activities to compromise genome integrity. Because the complexity of L1 biology is not understood, studies were conducted to evaluate the impact of L1 on epithelial to mesenchymal transition (EMT) in HepG2 cells. Forced expression of synthetic wild type or mutant (D702Y) L1 deficient in RT activity was associated with formation of cytoplasmic foci and mitotic aggregation with the nuclear compartment. Random de novo L1 retrotransposition was identified in cells expressing wild type L1, but not the D702Y mutant. Both synthetic wild type and mutant L1 induced marked reductions in the expression of epithelial markers and overexpression of mesenchymal markers. These data establish L1 as a key regulator of genome plasticity in HepG2 cells via RT-dependent and RT-independent mechanisms which do not couple to intrinsic retrotransposition activity.

Interoperability Case Study: Connecting Models across Exposure and Dose-Response Arenas for Characterization of Vinyl Chloride Hepatotoxicity.

The 2012 SOT workshop “Building for Better Decisions: Multi-scale Integration of Human Health and Environmental Data” advanced seamless data and model integration for support of sustainability and improved risk characterizations. It highlighted the need for interoperability of models across the source/exposure/dose-response/cost/benefit continuum. We conducted a case study to illustrate interoperability of a range of human health models and chemical-specific properties for comparison of exposures to vinyl chloride (VC). Exposure models used were the EPA's Assessment System for Population Exposure Nationwide (ASPN) using data of the National-scale Air Toxics Assessment (NATA) for inhalation, and the Human and Ecological Exposure and Risk in Multimedia Environmental System (HE2RMES) domain within the Framework for Risk Analysis in Multimedia Environmental Systems (FRAMES) with Data for Environmental Modeling (D4EM) of the Agency's integrated exposure modeling (iem) technologies. The dose-response was characterized by data on the Integrated Risk Information System (IRIS) and the physiologically-based pharmacokinetic (PBPK) model for VC available in the ATSDR PBPK toolkit. Output from ASPEN and HE2RMES for inhaled VC exposure illustrated the value of efficient model comparison if plug-and-play capability is provided by interoperability. Conflicting assumptions regarding key parameters between exposure and dose-response arenas were highlighted when computing aggregate risk from oral and inhalation routes; these conflicts must be rectified prior to proper model integration. This exercise readily shows how the ability to transparently integrate such models impacts current approaches and enhances understanding of each discipline. (The views expressed in this abstract are those of the authors and do not necessarily represent the views or policies of the U.S. Environmental Protection Agency or the Agency for Toxic Substances and Disease Registry.)

Abacuse Exposure Guideline Levels (AEGLs) are developed by the USEPA AEGL committee for controlling acute exposures to airborne chemical hazards. AEGLs are the threshold exposure limits for once-in-a-lifetime or rare chemical exposures at five exposure durations (1/6, 1/2, 1, 4, 8h). AEGLs are derived from various published and unpublished experimental studies described in the Technical Support Documents prepared by the committee. An AEGL concentration (C) for a specific duration (t) is extrapolated from available experimental data. Extrapolations are carried out using the exponential function, $C_n = C_0 \times t^k$, where $n$, the temporal scaling factor (TSF), is chemical-specific. Preferably, it is derived experimentally, but thus far experimental TSFs for few chemicals have been derived. For most of 272 chemicals on the AEGL list, TSFs are unknown. For these chemicals, the AEGL committee has adopted a rule by which $n = 1$ when extrapolating from short- to short-term durations, and $n = 3$, when extrapolating from long- to short-term durations. However, for many chemicals with unknown TSFs, this rule has been abandoned in favor of chemical-specific information deemed appropriate by the committee. Thus, the AEGL database contains rich expert-validated chemical-specific information about temporal extrapolation. The objective of the present study was to extract this information. Using regression analysis for each chemical in the database, a surrogate TSF was derived. In addition to being chemical-specific, TSFs were found to be health-effects-specific. The relationship between median TSFs for mild/reversible (n=2.9, AEGL-1 tier), disabling/irreversible (n=2.0, AEGL-2), and life-threatening effects (n=1.8, AEGL-3) was $n_2 > n_1 > n_3$. A geometric mean TSF for the AEGL-3 tier, $n_m = 1.88$ [95% CI: 1.77, 1.99], was closest to $n = 1$ in Haber’s rule, which applies to combat exposures to chemical warfare agents. This observed agreement suggests that model-derived TSFs may be appropriate for temporally extrapolating the endpoint of AEGL concentrations to durations of exposure for which experimental durations are not available.
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A Novel Screening System for Classifying Potential Health and Environmental Impacts Associated with Hydraulic Fracturing Fluids.

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Public opposition to hydraulic fracturing or “fracking” is being witnessed across many communities in the U.S.A., Canada and elsewhere. Concerns over this practice include the potential impacts of fracking on groundwater and surface water quality, air quality, human health, and terrestrial and aquatic ecosystems. These concerns have prompted some jurisdictions to place temporary moratoriums on the practice until a better understanding of these potential impacts is gained. Numerous initiatives are now underway to heighten this understanding, including measures aimed at increasing awareness of the chemical additives that are used during the process. Many jurisdictions are now mandating public disclosure of these additives, and chemical registries that list the additives delivered down-hole on a well-by-well basis have emerged. Although these registries identify the chemicals used, they provide no indication of the potential health and environmental risks that the additives could present. To help fill this void, a novel risk-based screening system was developed to improve the understanding of the potential health and environmental impacts of the chemical additives used in hydraulic fracturing. The system relies on the results of toxicity and environmental fate studies for a number of key endpoints to assess the potential risks involved. The endpoints include acute and chronic oral toxicity, reproductive and developmental toxicity, mutagenicity, carcinogenicity, aquatic toxicity, environmental persistence and bioaccumulation potential. The outcomes of the screening process are used to assign the chemical additive products to one of several categories depending upon the potential risk involved. This categorization process allows the product users to take appropriate steps to manage the potential hazards. The screening system has been used to classify more than 1,500 products to date, has proven to be reliable and effective, and is currently being adopted on an industry-wide basis in Canada.

Analysis of the Validated Epiderm Skin Corrosion Test (EpiDerm SCT) and a Prediction Model for Sub-Categorization According to the UN GHS and EU CLP.

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Skin corrosion refers to the production of irreversible damage to the skin manifested as visible necrosis through the epidermis and into the dermis. In 2004, OECD adopted two ECVM-validated reconstructed human skin model assays (EpiDerm and EPISKIN) for testing skin corrosion (OECD TG 431). However, OECD TG 431 does not satisfy international labeling guidelines for transport of dangerous goods since none of the methods were validated for sub-categorization. The UN-GHS utilizes 3 corrosion sub-categories (1A-very dangerous, 1B-medium danger and 1C-minor danger). Labeling a chemical as 1A has important consequences for transport, including very small volume package limits for air transport, dangerous goods for road and rail transport, and largeamy limits for sea and river transport. The EpiDerm SCT was validated for sub-categorization based on a 3 min endpoint into the validated EpiDerm SCT deficition of corrosives into sub-category 1A, 1B/1C, and NC. Adoption of the new Specificity for NC chemicals was 80%.

Provisional advisory levels (PALs) are tiered exposure limits for toxic chemicals in air and drinking water that are developed to assist in emergency responses. Physiologically-based pharmacokinetic (PBPK) modeling can support this process by enabling extrapolations across doses, and exposure routes, thereby addressing gaps in the available toxicity data. Here, we describe the development of a PBPK model for Fentanyl – a synthetic opioid used clinically for pain management – to support the establishment of PALs. Starting from an existing model for intravenous Fentanyl, we first optimized distribution and clearance parameters using several additional IV dataset. We then calibrated the model for oral and inhalation exposures using pharmacokinetic datasets for various Fentanyl formulations. Predictions of the calibrated model were in good agreement with the >50 datasets considered here. For aerosolized pulmonary Fentanyl, F=1 and c90·1<min indicating complete and rapid absorption. The F value ranged from 0.35 to 0.74 for oral and various transmucosal routes. Oral Fentanyl was absorbed the slowest (90 – 300 mins); the absorption of intranasal Fentanyl was relatively rapid (90 – 20-40 mins); and the various oral transmucosal routes had intermediate absorption rates (90 – 160-300 mins). Based on these results, for inhalation exposures, we assumed that all of the Fentanyl inhaled from the air during each breath directly, and instantaneously enters the arterial circulation. We present model predictions of Fentanyl blood concentrations in oral and inhalation scenarios relevant for PAL development, and provide an analytical expression that can be used to extrapolate between oral and inhalation routes for the derivation of PALs.


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Recent National Research Council recommendations on the process of hazard assessment include improving transparency and standardization of evidence collection, selection, presentation, and evaluation. Per this recommendation, we designed
a process that was tested using di(2-ethylhexyl)phthalate (DEHP) liver tumorigenesis as a model. In addition, relevant synonyms and wildcards for DEHP and its metabolites, MOAs, and species of interest were tested in PubMed. Specific tiered assessment criteria for exclusion/inclusion into the search were also defined and literature was searched and screened according to the tiered criteria. For MOAs with large databases, additional screening criteria were applied through manual curation of the articles. The evidence gathering process was documented as a literature tree, clearly describing each step of the search, inclusion/exclusion, and curation. Studies satisfying the assessment criteria were summarized into evidence tables. Narratives for each liver cancer related MOA were composed. Literature was organized in the EPA’s Health and Environmental Research Online (HERO) database using tags in accordance with assessment criteria. This approach streamlines and exemplifies transparent mechanistic evidence capture and reporting for DEHP liver carcinogenicity. Large amounts of information are effectively managed and appraised, and enabled for analysis and interpretation of cancer genome/cancer pharmacogenomics. This further supports the development of a comprehensive chemical risk assessment framework for prioritized chemicals as part of the National Environmental Research Online (HERO) database using tags in accordance with assessment criteria. This approach streamlines and exemplifies transparent mechanistic evidence capture and reporting for DEHP liver carcinogenicity. Large amounts of information are effectively managed and appraised, and enabled for analysis and interpretation of cancer genome/cancer pharmacogenomics. This further supports the development of a comprehensive chemical risk assessment framework for prioritized chemicals.

**712 Material Threat Assessment Chemical and Scenario Selection Process.**

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The Department of Homeland Security (DHS) Chemical Security Analysis Center (CSAC) and the Department of Health and Human Services (DHHS) Biomedical Advanced Research and Development Authority Division of Analytic Decision Support (BARDADS) jointly assess public health risks associated with potential chemical agent attacks on the U.S. homeland. To provide a realistic appraisal of the type of threats chemical agents pose to the U.S. populace, the CSAC and BARDADS evaluate a broad range of chemicals to assess an adversary’s potential to cause significant adverse consequences. This is accomplished through the development of Material Threat Assessments (MTAs), which examine the potential acquisition of a chemical and whether effective dissemination on a target can cause significant severe or lethal effects. Each MTA consists of an analysis of a single chemical or group of chemicals that have a similar mechanism of action, while modeling these chemicals across a standard set of scenarios to enable comparison among relevant criteria. Each MTA includes a comparison of public health consequences of a chemical attack with and without the use of consequence mitigating countermeasures. This presentation describes how the Chemical Terrorism Risk Assessment (CTRA) is used to inform the MTA process and describes the tools and assumptions employed to select high risk and high consequence chemicals and scenarios.

**713 Use of REACH Registration Data for Improving Thresholds of Toxicological Concern (TTC).**

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The threshold of Toxicological Concern (TTC) concept is useful to identify human exposures that are so low that in-depth toxicological investigations are expendable. This is called “exposure-based waiving”. Exposure-based waiving serves to focus available resources on substances with relevant human exposure potential. Important work into establishing TTC values has been published by Munro et al. (1996). The initial report used a database of 613 organic substances compiled from publicly available sources. In total, 2941 NOELs were collected in this fashion. The Munro concept used the Cramer classification to categorise substances according to their hazard potential. We broadened the TTC database by including NOAELs published on the ECHA website as per 19 July 2012, containing data for more than 7600 substances. Only non-gaseous mono-constituent substances with oral NOAELs were included in the TTC database. Organophosphates and genotoxic substances were excluded from the database as well as NOAELs obtained for surrogate substances. NOAELs for all systemic endpoints (general toxicity, developmental toxicity, fertility, neoplasia) were taken into account. Where appropriate, default assessment factors of up to 6 were used to establish chronic NOAELs for each substance. For every eligible substance, we collected the published CLP category for acute oral toxicity as a potential predictor of overall hazard potential. This gives rise to five categories of acute oral toxicity. A TTC is calculated from the 5th percentile of NOAELs in each of these categories using the REACH rules for establishing DNELs for workers and the general population. This poster presents the preliminary results for more than 1500 substances. The results indicate that the TTC concept becomes more robust when using the very broad ECHA database. It also suggests that acute oral toxicity categories can be used as a predictor for the overall hazard potential of a substance.

**714 Evaluation of the ToxRtool for Assessing Quality of Toxicological Studies for Risk Assessments.**

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In order to improve transparency and consistency, both the European Union and National Academy of Sciences have recommended the standardization and documentation of criteria used to evaluate the quality of studies considered in health risk assessments of xenobiotics. Although peer-reviewed publication is an important criteria for judging scientific data, the quality of studies and reporting of results vary tremendously. To find a means to efficiently assess the quality of these studies, we evaluated a publicly available algorithm (Toxicological data Reliability assessment Tool, ToxRtool) recently developed by the European Commission. The ToxRtool builds on the Klimisch score, a method for categorising the reliability of toxicological studies, but adds important criteria that should be present in the high-quality journal article used to support health risk assessments. To evaluate the ToxRtool, 20 peer-reviewed studies on thyroid disrupting chemicals were selected, ranging in quality from excellent to poor as determined by a priori by a thyroid expert on the team. Eight scientists with various levels of expertise then used the ToxRtool to evaluate the papers, and the consistency and reliability of scores were compared across evaluators. Scores were most consistent for previously judged ‘high-quality’ papers with greater variability among scores as the quality of papers diminished. The basis for inconsistencies appeared to be related to the subjective nature and lack of clarity of responding to many criteria. In conclusion, by providing more objective and standardized instructions for responding, the ToxRtool would likely be extremely useful for systematic evaluations of peer reviewed studies being considered for health risk assessments.
GO-Quant Systems-Based Quantitative Analysis of Dynamic Signaling Pathways during Neurodevelopment and Implication for Risk Assessment.

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The most critical biological process during neurodevelopment is the timely controlled signaling pathways. Many reports provided qualitative data demonstrating the critical role of neuro-developationally related pathways. However, only few have provided quantitative evaluation of the dynamic pathways during these processes. These differences in dynamic signaling pathways are believed to be tightly associated with the sensitivity of window of exposure to toxicants, but still have not been clearly examined. Genome-wide scale evaluation of gene expression has proven to be a powerful tool in understanding molecular mechanism during neurodevelopment. Through analysis of these genomic data during development, we inevitably lead to a greater understanding of how changes in transcriptome relate to functional and structural changes in the neurodevelopment. However, the integration of these gene expression data into a description of dynamic signaling pathways within specific events of development has proven to be a difficult task.

We constructed and established meta-database for neurodevelopment and applied systems-based GO-Quant program to quantify strains, could predict the dynamic stage-specific landmark of signaling pathways. Our GO-Quant analysis with rat gene expression array data demonstrated the dynamic changes of multiple signaling pathways during different stages of development. The mapping of the dynamic pathways greatly improves our understanding of time-dependent role of each signaling pathway during neurodevelopment. The systems-based identification of developmental stage-specific landmark signaling pathways helps to create sensitive, breakthrough early biomarkers for neurodevelopmental toxicity.

Safety Assessment of a Novel Antibiotic Using a Mouse Population-Based Approach Predicts Risk of DILI in Humans Where Classical Models Fail.

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Development of the macrolide antibiotic PF-04287881 was suspended following elevations in liver function tests observed in a Phase 1 clinical trial. The potential for drug-induced liver injury (DILI) due to this compound was not predicted by standard nonclinical toxicology studies. We hypothesized that a mouse diversity panel (MDP), comprised of genetically diverse inbred strains, could predict the DILI potential of PF-04287881. Additionally, using the MDP offers the ability to identify the critical role of neuro-developationally related pathways, but still have not been clearly examined. Genome-wide scale evaluation of gene expression has proven to be a powerful tool in understanding molecular mechanism during neurodevelopment. Through analysis of these genomic data during development, we inevitably lead to a greater understanding of how changes in transcriptome relate to functional and structural changes in the neurodevelopment. However, the integration of these gene expression data into a description of dynamic signaling pathways within specific events of development has proven to be a difficult task.

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An Investigation of Factors Associated with Mortality Patterns across the Midwestern United States.

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Various studies have indicated that mortality rates are positively correlated with social inequalities, air pollution, elevated ambient temperature, age, availability of medical care and other factors. This study develops a model that uses indicators for multiple factors to predict the mortality rates for selected diseases by county across the Midwestern United States. A total of 1,055 counties from 12 states in the Midwestern region were studied. Systematic random sampling was used to select a subset of counties from the 1,055 counties to validate the model. Appropriate statistical analyses were used to predict the relationship between environmental pollutants, socio-economic factors, risk factors, social capital, weather, crime, social and other factors to explain variations in county-specific mortality rates for the leading causes of death and lifespan in the US. It is anticipated that the study would suggest the complex inter-relationships of multiple factors that influence mortality and lifespan, and suggests the need for a better understanding of the pathways through which these factors, mortality, and lifespan are related at the community level, particularly in urban versus rural settings. The resulting findings might assist policy makers at local, state, and federal levels in developing effective prevention strategies in this region.
a. Integrated Mode-of-Action (MoA) and human relevance investigations – for example, anchoring in vitro MoA test concentrations to the in vivo effect levels in animals.

b. Setting of ‘internal’ Reference Doses (RIDs) – for example, proposing the corresponding blood concentration from critical in vivo studies used to set RIDs (such as the acceptable daily intake (ADI) or acute reference dose (ARID), as an ‘internal’ RID.

Traditional human health risk assessments include Uncertainty Factors (UFs) to account for toxicokinetic (TK) and toxicodynamic differences between animals and humans. Predictions of systemic exposure in humans, based on animal TK data or measured human biomarker levels, could allow for science-based refinements in the UFs related to interspecies differences in TK. These refinements could provide more realistic reference dose values, while predictions of human blood levels could help to inform calculations of biomonitoring equivalents.

The aims of this project were twofold. The first was to develop a user-friendly modeling method that could accurately predict systemic bioavailability via parenteral exposure routes (i.e, dermal) in humans. Ideally, this model could predict exposure levels in both urine and blood. The second was to utilize animal TK data (measured or modeled from alternate exposure routes) to predict systemic exposure in humans from either occupational, bystander or residential exposures, using a simple TK modeling approach.

TK MOD® is an Excel-based program developed for prediction of blood levels of a test material and optimal sampling time selection for TK analysis. This program was successfully used to predict human blood levels, both, after exposure by parenteral routes (dermal), and using data derived only from animals. Three data-rich Dow AgroSciences molecules were used as case studies: haloxyfop, triclopyr and picloram. Attempts were made to achieve both of the above aims with each molecule, though this was not always possible due to limited available data. Overall, accurate predictions of human blood levels were achieved in most cases where adequate data were available. The only exceptions were due to large species differences between animal and human kinetics. A greater understanding of these animal/human system exposure comparisons will help to reduce interspecies uncertainty related to TK that are used in regulatory risk assessments.

The Chemical Terrorism Risk Assessment (CTRA) is a probabilistic risk assessment that considers threat, vulnerability, consequences, and mitigation to yield a comprehensive estimate of chemical terrorist risk to the Nation. The Medical Mitigation (MedMit) model is a key component of the CTRA; it is a stock and flow model with parameters defined by toxicology and emergency medicine subject matter experts. The MedMit model estimates the number and type of injured victims according to dose, and determines the number of victims that may be saved or benefitted by the public health response. This presentation will discuss the approach to parameter quantification. It includes the definition of chemical-specific toxicity values for life-threatening and severe injuries, and mild to moderate injuries, types, and the desire and benefits of including these different victim types in the Medical Mitigation model, the segregation of more than 100 chemicals into toxidromes, and the data collection process. The application of toxicology to the CTRA enables a more realistic representation of event consequences and the public health response. Consequently, the model makes it possible to assess the effectiveness of the existing public health response system and examine improvement strategies. Such a capability permits policy makers to make informed decisions regarding resource allocation, and assists responders’ understanding of chemical terrorist events and areas of potential improvement.

The h-NAG-1 Transgenic Mouse Serves As an Animal Model for Obesity.

H. Carlson-Lynch1, J. Stickleyn1, A. Bacon1, G. Diamond1, B. Thayer1, P. McClure1, L. Flowers2, C. Chen3, M. Gebhards4 and M. Pratt5. 1SRC, Inc., Annandale, VA. 2NIH, Research Triangle Park, NC.

NSAIDS-activated gene-1 (h-NAG-1) transgenic mice (C57BL/6-Tg(CAG-GDF15) over express human NAG-1/ Growth/differentiation factor-15 (GDF15). Hemizygous h-NAG-1 mice are significantly leaner than their wild-type littermates (GDF15) over express human NAG-1/ Growth/differentiation factor-15 (GDF15). Hemizygous h-NAG-1 mice are significantly leaner than their wild-type littermates. (h-NAG-1) transgenic mice (C57BL/6-Tg(CAG-GDF15)) were significantly decreased in h-NAG-1 mice on both diets. 

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The h-NAG-1 transgenic mouse (C57BL/6-Tg(CAG-GDF15) over express human NAG-1/Growth/differentiation factor-15 (GDF15). Hemizygous h-NAG-1 mice are significantly leaner than their wild-type littermates and have less abdominal white fat despite comparable food intake. To improve our understanding of the mechanisms of obesity development, h-NAG-1 mice and WT were put on a 10 or 60% diet for 12 weeks and blood was collected. Final body and abdominal white fat weights were increased in WT relative to h-NAG-1 mice. The h-NAG-1 mice on the high and low fat diet had lower circulating levels of the inflammatory adipokine leptin, although the difference on the high fat diet was much greater. The h-NAG-1 mice (F5.4 +/- 0.7 mg/ml) on the high fat diet (2.5 +/- 0.7; h-NAG-1 and 15.8 +/- 6.8 ng/ml; WT). Similarly, serum insulin levels were significantly decreased in h-NAG-1 mice on both diets. Because obesity, via

The Association of CTR and MM in a Map of Specificity Factors. P. Nance1, L. Whitmire2, J. A. Cox2. 1SRC, Inc., Durham, NC; 2NIEHS, Research Triangle Park, NC.

The association of CTR and MM in a map of specificity factors. A greater understanding of these animal/human system exposure comparisons will help to reduce interspecies uncertainty related to TK that are used in regulatory risk assessments.

An Approach for Characterizing Uncertainty in Relative Potency Factors.

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Relative potency factors (RPFs) for cancer may be calculated as the ratio of the slopes of two dose-response curves, each estimated by linear extrapolation from a benchmark dose (BMD) value. BMD values are subject to uncertainties stemming from study design features such as dosing interval and numbers of animals tested, as well as uncertainties pertaining to modeling, such as model choice, goodness-of-fit, and choice of benchmark response (BMR) relative to the observable data range. Quantitative characterization of the uncertainty in a RPF estimate is desirable both for comparing different estimates and for use in characterizing uncertainty in risk estimates obtained with RPF. Because it is estimated as a ratio, a RPF poses certain challenges with respect to statistical approaches to characterizing uncertainty. Analytic approaches to calculating a confidence interval for a ratio (e.g., Hinkley 1969) were explored as a means of characterizing the uncertainty in a number of RPF estimates for benzo[a]fluoranthene (relative to benzo[a]pyrene). In addition, a Monte Carlo simulation method was investigated. For both approaches, the RPF estimate was simplified as a ratio of BMDs estimated at the same BMR (10% extra risk), and the standard error of the BMD was approximated using the upper and lower 95% confidence limits reported by the BMD software. For the Monte Carlo method, the benchmark dose was simulated as a lognormal distribution. Each iteration of the Monte Carlo simulation consisted of random samples from the probability distributions of BMDs for benzo[a]fluoranthene and benzo[a]pyrene, and calculation of an RPF for each random draw. The Monte Carlo simulation approach, unlike the analytic approach, provided plausible confidence limits and therefore, useful quantitative estimates of the uncertainty in the RPFs. (The views expressed are those of the authors and do not necessarily reflect the views or policies of the U.S. EPA.)
higher leptin levels, can increase inflammatory responses, hNAG-1 mice and WT were injected with the inflammatory agent lipopolysaccharide (LPS) and serum inflammatory cytokine measured. The levels of KC, IL-6, MCP-1 and TNFα were lower in the leaner hNAG-1 transgenic mouse. The data indicate the hNAG-1 mouse may serve as a model for investigating obesity and how environmental exposures may be altered by obesity.

A possible association between use of the anti-diabetic drug exenatide and acute pancreatitis has been suggested by the analysis of post-approval adverse event data but remains controversial. Studies were undertaken to determine whether an animal model could be developed to reveal potential safety signals consistent with pancreatitis following exposure to drug. Subcutaneous exenatide injection (3, 10, or 30 μg/kg for 6 weeks) of Sprague-Dawley rats, caerulein-treated rats, and Zucker fatty diabetic rats demonstrated no consistent histopathologic pancreatic injury signal on standard chow or a high fat, high carbohydrate diet (HFD). However, C57BL6 mice placed on HFD for 6 weeks exhibited a dose and time dependent pancreatic responses to single daily subcutaneous exenatide injections (3, 10, or 30 μg/kg, for up to 12 weeks) that were exacerbated by the HFD. Focal areas of acinar cell necrosis progressing to edema, inflammation, and general tissue fibrosis and atrophy were identified in control mice but were much more prevalent in exenatide treated mice. Other findings included acinar cell hypertrophy and hyperplasia with increased incidence of autophagy and apoptosis in proportion to necrosis and inflammation. Regardless of exenatide exposure, mice on HFD had increased body weight and significantly elevated blood glucose and serum pro-inflammatory cytokines compared to mice on standard chow. The type of stress and injury produced in this study do not approximate the acute necrotizing pancreatitis described in adverse event reports but represent a reproducible pancreatic signal that might be useful for the preclinical safety evaluation of pancreatic injury induced by anti-diabetic drugs.

Extended Exenatide Treatment Causes Pancreatic Stress and Injury in a Rodent Model of Insulin Resistance.


Development of an Oral Exposure Model to Predict Drug Hypersensitivity Using Brown Norway Rat.


Behavioral, Neurochemical and Histological Characterization of a Novel MitoPark Mouse Model for Neurotoxicity and Neuroprotection Studies.


Parkinson’s disease is now recognized as a neurodegenerative condition caused by a complex interplay of genetic and environmental factors. The animal models presently available, created by neurotoxic insults or genetic defects, do not fully recapitulate the chronic and progressive nature of the nigrostriatal dopaminergic neurodegenerative process. Recently, the MitoPark mouse model was created by inactivation of mitochondrially transcription factor Tfam in the nigrostriatal pathway through the control of a DAT promoter. In this study, we tested the utility of MitoPark mice for neurotoxicological and neuroprotective studies using behavioral, neurochemical and histological analyses. Measurement of locomotor activity in MitoPark mice over 8-25 weeks revealed that motor function started to decline at 12 weeks of age and progressed over time, leading to severe deficits at 17-25 weeks.

728 Development of an Oral Exposure Model to Predict Drug Hypersensitivity Using Brown Norway Rat.

729 Behavioral, Neurochemical and Histological Characterization of a Novel MitoPark Mouse Model for Neurotoxicity and Neuroprotection Studies.

725 Extended Exenatide Treatment Causes Pancreatic Stress and Injury in a Rodent Model of Insulin Resistance.


726 Adipose Deficiency of Nrf2 in Ob/Ob Mice Results in Severe Metabolic Syndrome.


Nuclear factor E2-related factor 2 (Nrf2) is a transcription factor that functions as a master regulator of the cellular adaptive response to oxidative stress. Our previous studies showed that Nrf2 plays a critical role in adipogenesis by regulating expression of CCAAT/enhancer binding protein β and peroxisome proliferator-activated receptor γ. To determine the role of Nrf2 in the development of obesity and associated metabolic disorders, the incidence of metabolic syndrome was assessed in C57BL/6 mice with Nrf2 deficiency. Absence of Nrf2 in WAT resulted in an even more severe metabolic syndrome, the incidence of metabolic syndrome was assessed in C57BL/6 mice with Nrf2 deficiency. Absence of Nrf2 in WAT resulted in an even more severe metabolic syndrome.
MitoPark mice also showed a dramatic loss of striatal dopamine and its metabolites compared to wild type C57 mice. In order to determine the gene-environmental interaction, we exposed MitoPark mice to Mn (10 mg/kg, p.o) daily for 30 days. Motor deficits were significantly exacerbated in Mn-treated MitoPark mice beginning at week 10 compared to vehicle treated MitoPark mice. The depletion of striatal dopamine, DOPAC and HVA content in MitoPark mice was exacerbated in Mn-treated MitoPark mice. We also evaluated the neuroprotective efficacy of a novel mitochondrial-targeted antioxidant, mitoapocynin. Oral administration of mitoapocynin (10 mg/kg, p.o) three times a week to MitoPark mice between 13-24 weeks of age restored behavioral deficits, striatal dopamine depletion and TH neuronal cell loss in MitoPark mice. Our results demonstrate that the MitoPark mouse is an excellent model to study the gene-environmental interactions associated with mitochondrial defects in the nigral dopaminergic system as well as to evaluate the neuroprotective efficacy of novel neuroprotective agents.

### 730 Continuous Infusion in Genetically-Altered Down Syndrome Mice.

H. van Wijk, Covance Laboratories Ltd., Harrogate, United Kingdom. Sponsor: S. Kirk.

Performing femoral vein catheterisation surgery with tail cuff exteriorisation directly at the performing laboratory offers various advantages. Implanted mice avoid transport from a third party supplier post-surgery. This reduces the risks associated with post-operative transport (expression of disease), requires a shorter total recovery time, and allows utilisation of unusual strains not surgically available. In this study we investigated the feasibility of performing femoral vein catheterisation with tail cuff exteriorisation surgery in-house on B6EiC3Sn a/A-Ts(1716)65Dn (T65DN) Euploid T65Dn mice (Jackson Laboratories, Bar Harbour, Maine, USA) to avoid tube restraint necessary for iv bolus administration and allow behavioural monitoring. T65DN is a widely accepted mutant strain model of Down's syndrome allowing assessment of medicines for potential unwanted side effects in disease patients. The chromosomal complement of the T65Dn mouse is incomplete relative to conventional mice; however the animals do not require special husbandry conditions, housing or maintenance and are considered to show normal behaviour in most respects. They are not affected by heart abnormalities common in humans affected by Down's Syndrome.

The mean body weight of the control euploid mice is greater than the T65Dn mice (~40g and 30g respectively). The mean body length for T65Dn mice is shorter (~1 cm nose-base of tail: 1.5 cm nose-tip of tail) presenting challenges in surgery and tail cuff attachment. To ensure tail size was not likely to impede exteriorisation of the catheter, a range of tail cuff sizes were used. Of the 16 T65Dn and 16 wild type mice that were surgically prepared, only one T65Dn mouse failed to recover and was humanely killed three days post surgery. No unexpected clinical signs were seen after surgery and infusion and body weights started to regain six days post surgery, which is quicker than previously described by Arts et al. 2012.

In conclusion, in-house catheterisation of T65Dn mice was shown to be more humane than obtaining and transporting surgically prepared mice from a third party, reducing the chance of disease expression.

### 731 Generation and Characterization of a Humanized C. elegans Transgenic Animal That Results in Early Onset, Age-Dependent Complete Loss of DA Neurons.

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Background: Parkinson's disease (PD) is a slowly progressive neurodegenerative disease characterized by the selective loss of dopamine (DA) neurons. Despite over 50 years of intensive research into the disorder, the origin of the pathogenesis and the molecular determinants involved in PD have not been elucidated. A significant hindrance in dissecting these molecular components is the lack of facile in vivo genetic models to explore the mechanisms involved in age-dependent cell death. Aims/Objectives: Our goal in this study was to develop and characterize a viable C. elegans animal that has early onset, age-dependent complete loss of DA neurons. Methods: We utilized transgenic C. elegans, reverse genetics, biochemical assays, immunofluorescence, mRNA and miRNA arrays, RT-PCR, Western analysis, and neuronal morphology analysis to characterize expression, localization and the role of human alpha-synuclein, parkin and mutant genes in DA neuron patholgy. Results: We generated genetic crosses with C. elegans containing human A53T alpha-synuclein, an endogenous parkin mutation and other genetic modifications. One product of this study is the generation of a C. elegans transgenic animal that results in a robust DA neurodegeneration phenotype. Immediately after hatching, the animals have their full complement of DA neurons as determined by confocal analysis and DA levels by HPLC, yet animals approximately 8 days old exhibit a complete loss of DA neurons. Our data suggests that post-translational modifications play a significant role in PD-associated genes and toxicants. Conclusions: We have generated a novel transgenic, the first animal (vertebrate or invertebrate) that has complete loss of DA neurons as a function of age. This animal is proving to be invaluable in determining molecular basis of gene- and toxin- associated DA neuron vulnerability. Support: NIEHS ES014459 and ES003299 to RN, and EPA STAR Graduate Fellowship to NVD.

### 732 Development of Animal Models of Idiosyncratic Drug-Induced Liver Injury.

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Background: Idiosyncratic drug-induced liver injury (IDILI) remains a serious health problem and it also significantly increases the risk of drug development. Unfortunately, not much is known about the mechanism of IDILI and a main reason is because there are no good animal models. We set out to develop animal models of IDILI by focusing on two drugs: Isoniazid (INH) and Amodiaquine (AQ), both of which cause IDILI in humans. Results: INH was found to covalently bind to mouse and to a lesser extent rat liver macromolecules. Treatment of Balb C, C57BL/6 mice, Wistar or Brown Norway rats with INH did not result in significant liver injury. Treatment of C6h-b+/-, PD1/- and NAT 1/2/- mice resulted in mild liver injury. One possible reason that these animals did not develop more extensive liver damage is immune tolerance. Immunization of C57BL/6 mice with S100 hepatic protein modified with INH produced a greater degree of autoimmune hepatitis than S100 alone, but when INH was subsequently given orally this prevented the autoimmune hepatitis. Treatment of female C57BL/6 mice with AQ resulted in a mild transaminase (ALT) increase at week 3 and this resolved by week 5. Immunohistochemical staining and flow cytometry showed that AQ-induced liver injury in C57BL/6 mice was associated with infiltration of immune cells such as F4/80, CD11b, CD4, CD8 and CD45R in the liver and spleen. Treatment of C6h-b+/- and PD1/-/- mice with AQ resulted in more severe liver injury, but ALT still resolved despite continued treatment. Treatment of C6h-b+/- mice with FICZ and Anti-CD25 Ab to break immune tolerance was not effective. Treatment of RAG1/-/- mice also resulted in a mild increase in ALT but the ALT did not appear to resolve as it did in the C57BL/6 mice. Conclusion: These data suggest that the dominant response to these drugs is immune tolerance. Treatment of female C57BL/6 mice with AQ results in delayed onset of mild liver injury which resolved despite continued treatment; this is similar to what happens in humans and this model is being further characterised. Supported by grants from CHHR.

### 733 Role of Myeloid-Derived Suppressor Cells in a Murine Model of Drug-Induced Liver Injury Mediated by the Adaptive Immune System.

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Although clinical evidence suggests that many cases of serious idiosyncratic drug-induced liver injury (SIDILI) are mediated by hepatic protein adducts of drugs and the adaptive immune system, detailed experimental proof for this mechanism of toxicity has remained elusive due to the lack of animal models. We have hypothesized that SIDILI is as rare in animals as it is in humans due at least in part to the tolerogenic nature of the liver, which consists of multiple negative regulators of the adaptive immune system. This idea has now been tested in an established murine model of halothane-induced liver injury where the toxicity is initiated by the metabolism of halothane to form trifluoroacetylated liver proteins and enhanced by the innate immune system, which are prerequisites for activating the adaptive immune system. Twenty-four hours after female Balb/CJ mice were treated with halothane, analysis of the liver revealed perivenous necrosis and an infiltration of CD11b+ Gr-1High neutrophils, as reported by other researchers. Further study revealed that the neutrophils contained a subpopulation of myeloid-derived suppressor cells (MDSC) that inhibited the proliferation of both CD4+ and CD8+ T cells isolated from naive mice. When MDSC were depleted from the liver of Gr-1+ mice on two treatments with halothane, enhanced liver injury was observed nine days after the second exposure of halothane as compared to mice that were pretreated with isotype control antibodies before halothane treatments. Moreover, the liver injury was associated with elevated levels of hepatic T cells and serum antibodies that both reacted with trifluoroacetylated liver proteins isolated from halothane treated mice. Collectively, our data provides a rational approach for developing animal models of SIDILI mediated by the adaptive immune system and suggests that deficiencies in liver tolerance may predispose patients to SIDILI.
Defective Platelet Function in a Mouse Model of Progressive Cholestasis.

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We have recently demonstrated that Abcb11-deficient mice recapitulate the human genetic disease PFIIC2 (progressive familial intrahepatic cholestasis type 2) attributed to loss of function of the ABC transporter, ABCB11 (1). Cholestatic liver disease leads to coagulopathies, however it is unclear if these are due to defects in the liver’s production of fibrinogen and procoagulant factors or defects in platelet function. By metabolomic analysis we demonstrated that lysophosphatidylcholine (lysoPC) (16:1 and 18:1) concentrations were elevated 2 to 5 times (vs. WT) in the liver and sera of Abcb11-null mice before evidence of frank liver damage. Because lysoPC inhibits platelet aggregation (2) we hypothesized that platelets of Abcb11-null mice might have aggregation defects due to elevated serum lysoPC. Platelet counts were comparable between the Abcb11-null mice and WT mice and platelet-rich plasma was used to test platelet aggregation. Agonist-induced aggregation of Abcb11-null platelets was impaired for agonists: ADP, collagen and thrombin. This ubiquitous defect is consistent with reduced expression of dense granule marker P-selectin and platelet receptor protein GPVI. As there are no changes in the downstream signaling genes P2y1, Syk, and FcYr1, in platelets of abcb11-null mice; our results suggest that defective aggregation of Abcb11-null platelets is the result of down-regulation of GPVI and P-selectin which may be secondary to lysoPC exposure. 1. Zhang Y, Li F, Patterson AD, Wang Y, Krausz KW, Neale G, Thomas S, Nachagari D, Vogel P, Vore M, Gonzalez FJ, and Schuetz JD. (2012) Abcb11 Deficiency Induces. Cholestasis Coupled to Impaired beta-Fatty Acid Oxidation in Mice. J Biol Chem 287, 24784-94. 2. Yuan Y, Jackson SP, Newham HH, Mitchell CA, Salem HH. (1995) An essential role for lysophosphatidylcholine in the inhibition of platelet aggregation by secretory phospholipase A2. Blood 86, 4166-74.

Validation of ROS As a Toxicity Marker in Zebrafish.

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Oxidative stress, associated with an increased level of reactive oxygen species (ROS), is a key factor in both drug-induced toxicity and disease pathogenesis. However, conventional methods for assessing ROS damage are labor intensive and slow hampering widespread use as a toxicity marker. Because of their genetic and physiological similarity to humans, zebrafish have shown promise as an efficient and predictive animal model for assessing drug toxicity, safety and efficacy. Similar to effects in mammals, ROS levels have been shown to increase in zebrafish after drug treatment and irradiation. In this study, we developed a quantitative whole zebrafish microplate-based ROS assay format that relies on a commercially available fluorogenic dye (4-[and]-6)-chloromethyl-2,7-dichlorodihydrofluorescein diacetate, acetyl ester, CM-H2DCFDA). We validated the assay using 7 characterized mammalian ROS inducers: TSA [Trichostatin A], PMA [4]-phorbol 12-myristate 13-acetate, Cisplatin, DCA [Dichloroacetate], Menadione, TBHP [tert-Butyl hydroperoxide] and Ethanol and 1 negative control compound, NAC [N-Acetyl-L-cysteine]. In order to optimize the assay, we assessed ROS level by zebrafish developmental stage, linear relationship between number of animals per microwell and fluorescence intensity, signal to noise (S/N) ratio, effect of carrier DMSO, assay specificity, and assay reproducibility and robustness.

To confirm results using the whole animal microplate format, we also visually assessed site of ROS induction in transparent animals using fluorescent based morphometric image analysis. Using both the microplate format and whole mount morphometric image analysis, compared to mammals, the overall correct prediction rate in zebrafish was 100%, which according to the European Centre for the Validation of Alternative Methods (ECVAM), is considered "excellent". These results underscore the high conservation of toxicity pathways among species and support use of the whole zebrafish ROS assay as a rapid, predictive in vivo assay for compound screening.

The Effect of Fluoride on the Bones and Teeth of ICR-Derived Gromerulonephritis (ICGN) Mice by Subacute Exposure.

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Dental fluorosis and osteofluorosis by drinking water contaminated with fluoride (F) have been reported in China and India. Because fluoride is excreted from the kidney, the toxic effects of F may be affected more significantly by F in ICR-derived gromerulonephritis (ICGN) mice that have commonly been used as a model of renal failure. In this study, we administered F to ICGN and ICR mice to compare the effects of F on the teeth and bones between mice with and without renal failure. Both male and female ICGN and ICR mice were administered F at the concentrations of 0, 25, 50, 100, and 150 ppm in the drinking water for 4 weeks. The mice’s femurs were sampled and photographed with a standard aluminum scale. The photographs were analyzed by microdensitometry, and the indexes calculated were: bone mineral content and bone density. The teeth were decalcified in EDTA solution and stained with HE. All the ICGN mice exposed to 150 ppm died. However, no ICR mice died. The mean bone density was significantly lower in the female ICGN mice exposed to 150 ppm compared to the control. The mean mineral content and bone density of the female 100 ppm ICGN group were significantly lower than those of the control. There were no significant differences in the bone indexes among the male ICGN groups. In ICR mice, the mean bone mineral content and bone density were higher in the 150-ppm group for both male and female. The heterogeneity of calcification in the tooth enamel was observed in the ICGN mice exposed to 150 ppm. The decrease in bone density in the mice exposed to F may be due to systemic effects of F. ICR mice exposed to F may be used as a model of osteofluorosis.

A Population-Level Mouse Model to Investigate the Genetic Determinants of Susceptibility to Environmental Toxics.

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Carcinogenesis biosays have lacked the genetic component critical for the evaluation of underlying disease susceptibility genes. Here we used a systems biology approach to model the genetic heterogeneity of exposed human populations, including the inter-individual variation in drug metabolism and transport. We developed a novel intercross population derived from FVB/N(ABcb1a/b-/-) x CAST/EiJ F3, a multi-drug resistant p-glycoprotein knockout mouse model, and CAST/EiJ a wild-derived strain that is genetically distinct from the FVB/N background. Using this intercross to model human biology, including using a western diet, environmentally-relevant doses of trichloroethylene (TCE) and inorganic arsenic (iAs), were evaluated for their effects on toxicity susceptibility. A study cohort of 900 FVB/N(ABcb1a/b-/-) x CAST/EiJ F3 mice was divided into nine dose groups, each containing 50 female and 50 males, and was administered TCE and sodium arsenite via the drinking water and chow, respectively, at environmentally-relevant concentrations for 56 weeks using a dose-ratio approach. At harvest, biofluids and tissues were either formalin fixed for pathology, or flash frozen for molecular analysis. Mice co-exposed to TCE and iAs developed hepatocellular carcinoma, as well as renal cysts, clear cell foci and dilated renal tubules consistent with human renal disease pathogenesis. Whole genome expression analysis of kidney tissues revealed a distinct response signature for the combinatorial exposures, including overexpression of oncogenes implicated in the development of human renal cell carcinoma. Genome-wide alleleotyping of each mouse using a 7,851 SNP genotyping array will reveal the genetic variants underlying toxicity response. This experimental paradigm has successfully modeled the population-level variability in disease response and has the potential to identify individuals sensitive to toxicity.
Human Antibodies Can Cross Guinea Pig Placenta and Bind Its Neonatal Fe Receptor: Implications for Evaluating Immune Therapy during Pregnancy.

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Disclaimer: The findings and conclusions in this presentation have not been formally disseminated by the Food and Drug Administration and should not be construed to represent any Agency determination or policy.

Despite increased use of monoclonal and polyclonal antibody therapies, including during pregnancy, there is little data on appropriate animal models that could humanly be used to understand determinants of protection, and to evaluate safety of these biologics in the mother and the developing fetus. We have demonstrated that pregnant guinea pigs can transport human IgG tranplacentally at the end of pregnancy. Using an intravenous anti-hepatitis B specific immune globulin preparation as an example, we show that IgG subclasses one, two and three are measurable in both the sows and their piglets. Given that IgG4 can be transferred tranplacentally in the guinea pig, our observations indicate that all human IgG subclasses can pass guinea pig placentas during the last third of pregnancy. In addition, we found that all human IgG subclasses are capable of binding to a soluble form of the guinea pig neonatal Fe receptor in vitro in a manner similar to that demonstrated for the human variant. This result suggests that tranplacental transport of human IgG subclasses in guinea pig mirrors the receptor-based mechanism seen in humans. Together, our studies lay the groundwork in introducing pregnant guinea pigs as an appropriate model for the evaluation of antibody therapies during pregnancy and advancing the health of women and neonates.

Development of a Phosgene-Induced Acute Lung Injury in the Conscious Pig.

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The toxic industrial chemical (TIC) phosgene (CG) is a reactive intermediate used in a range of industrial processes. Exposure to high concentrations of CG results in an asymptomatic period (2-6 h) before pulmonary oedema develops. There are no specific medical countermeasures for such poisoning, treatment being supportive in an intensive care setting. Evidence based treatment guidelines are needed for health care practitioners to guide treatment. Small animal models have been used to screen candidate therapies, extrapolation of therapeutic benefit to man requires verification in a larger animal model e.g. the pig. There is, therefore, a requirement to develop a conscious large animal model of CG-induced lung injury to verify efficacious treatments. If successful, this model will allow assessment of therapeutic interventions to CG-induced lung injury in a species closer to man.

After a period of socialisation, animals were surgically prepared to allow physiological measurements (arterial blood pressures, ECG, temperature and activity via telemetric implan) and blood samples (via exteriorised catheters) to be taken. After a 1 week recovery from surgery and baseline measurements (1 week) animals were exposed to either air or phosgene under anaesthesia, then recovered and monitored for 24 h.

Air-exposed and phosgene-exposed animals were successfully recovered and monitored to 24 h post exposure. The phosgene-exposed animals had a severe but non-lethal phosgene-induced lung injury at 24 h compared to air-exposed controls, demonstrated by changes in arterial blood gas measures, gross pathology and histopathology.

The feasibility of this unique model has been demonstrated. Further refinement of the telemetry technology may be required to add robustness and improve confidence in extrapolation to man. We intend to use the model to test therapeutic candidates proven efficacious in a small animal model.

Sex Differences in Inflammatory Response in a DAPM-Induced Rat Model of Pulmonary Hypertension.


Pulmonary Arterial Hypertension (PAH) is a cardiovascular disorder characterized by elevated pulmonary artery pressure as a result of arterial wall thickening. Patients survive, on average, 2.8 y after diagnosis and are 3-4 times more likely to be women than men. There is no cure - the few therapies available can only slow progression. Our purpose was to develop a relevant animal model of PAH in order to identify sex differences that contribute to disease progression, and in doing so, identify potential new therapeutic strategies. 4,4′-Methylenedianiline (DAPM) is an aromatic amine used industrially in the synthesis of polyurethanes. Chronic, intermittent treatment of rats with DAPM results in medial hypertrophy of pulmonary arteries, exclusively in females, coupled to increases in pulmonary arterial pressures. After 12 wk, significant increases in plasma levels of endothelin-1 (ET-1) and serotonin (5-HT), but decreases in nitrite (NO2−), were observed in females (not males) treated with DAPM. A decrease was observed in the serum ratio of the estrogen metabolites 2-hydroxyestradiol (2-OHE) and 2-hydroxyestrone (2-OHE; p < .07). In females, ET-1, NO2− and 2-OHE/2-OHE were significantly correlated with peak pressure gradient, an indirect measure of pulmonary arterial pressure. In DAPM-treated females, serum IL-10 decreased 53% (p = .0129). INF-γ increased by 48% and IL-6 increased 3-fold (as determined by Bio-Plex cytokine assay). These changes were not significant. However, there were significant interactions between sex and treatment. Future studies will address the contributions of these inflammatory cytokines in PAH initiation and progression in females.
Bleomycin Sulfate-Pulmonary Fibrosis has been extensively described in mice. However, to our knowledge few non-murine animal models have been implemented before. Here we investigated the pathogenic/inflammatory potential of Bleomycin administration in Cynomolgus Macaques (Macaca fascicularis), a surrogate model for human pulmonary fibrosis. Fifteen Cynomolgus Macaques (6 inulin resistant on high fat diet and 9 on standard primate chow) were challenged via a bolus insufflator with varying doses of Bleomycin Sulfate (0.1-1.9 mg/kg). Animals were followed during 10 weeks of study by twice daily cage-side clinical observations. Bronchoalveolar lavage (BAL) and pulmonary function tests (PFTs) were performed biweekly until necropsy. PFTs were completed for each animal (previously anesthetized) using a whole body plethysmograph. A subset of inflammasome potential markers was further analyzed in BAL cell mRNA. Macrophage lung abnormalities were examined at necropsy. Increases in PYCARD and TXNIP were noted in all animals that received high fat diet compared to those on normal primate chow irrespective of the bleomycin dose. Interestingly, NLRP3, PYCARD, PRX2X7, TXNIP, and IL-18 mRNA expression showed an inverse relationship with the bleomycin dose for both feed types. The development of fibrosis in monkeys mimics more of a chronic process as seen in humans. Though our animals presented structural/molecular changes, all of them were asymptomatic. Animals fed a high fat diet have increased inflammasome activation than those fed the standard diet however there is not a consistent association between the bleomycin dose and inflammasome activation. These findings suggest that Cynomolgus Macaques may be a potential animal model for studies of Bleomycin-induced pulmonary fibrosis, and this might prove to be a prerequisite for translational research.

Cynomolgus Macaque (Macaca fascicularis) As a Potential Animal Model for Studying Bleomycin-Induced Pulmonary Fibrosis. M. Doyle-Eisele1, J. Villallba1, Y. Shi1, J. D. McDonald1 and I. O. Rosas1,2. 1ERRR, Albuquerque, NM; 2Harvard Brigham and Women’s Hospital, Boston, MA.

Hepatic biotransformation is an important determinant of chemical bioaccumulation. Consequently, inappropriate use of in vitro models can be made using estimates of chemical biotransformation rates. Cryopreserved trout hepatocytes have previously been used to measure the clearance rates of some compounds. The use of this information within a regulatory context requires, however, that such results be reproducible across laboratories. In this study, three independent laboratories performed a round-robin study using cryopreserved rainbow trout (Oncorhynchus mykiss) hepatocytes. Six compounds were selected for testing: benzo[a]pyrene, 4-nonylphenol, di-tert butyl phenol, fenthion, methoxychlor and o-terphenyl. Pre-trial studies were performed to streamline the assay protocol, standardize hepatocyte counting procedures, and characterize potential sources of variability. The results confirmed first-order depletion kinetics for the selected compounds and highlighted the effects of assay temperature as well as lot variability. Each laboratory then conducted clearance assays for the six compounds using a substrate depletion approach. The analyses for each substrate were conducted at one institute to focus the comparison on the assay itself. Compounds determined to be poorly (o-terphenyl; <0.05 ml/100 mg/mL) or rapidly metabolized (benzo[a]pyrene; >0.3 mg/mL/100 mg/mL) were similarly determined across laboratories. Coefficients of variation across the three laboratories were generally 30% or better, suggesting this method of determining intrinsic clearance is transferable and reproducible. This inter-laboratory comparison strongly supports the use of cryopreserved trout hepatocytes as a tool for estimating hepatic clearance for bioconcentration factor prediction.


Dichloroacetate (DCA) is an investigational drug for mitochondrial diseases, pulmonary arterial hypertension (PAH), and cancer in humans. It is metabolized to glyoxyacetate by glutathione transferase zeta 1/malelacteocactase isomerase (GSTZ1/MAAI), which also catalyzes the penultimate step in tyrosine catabolism. DCA is a mechanism-based inhibitor of GSTZ1/MAAI, thereby inhibiting its metabolism and causing accumulation of reactive tyrosine intermediates, such as malonylacetate (MA). Malepate transformation in GSTZ1 also confers variable rates of DCA biotransformation that may impact chronic safety. Dogs express GSTZ1/MAAI and develop PAH and cancer; thus, they may be a valuable model of human diseases in which DCA is used. Accordingly, we characterized the pharmacokinetics and dynamics of DCA in a canine model.

Early Juvenile Exposure to High-Dose Acetaminophen Impacts Adult Murine Social Behavior in a Strain-Dependent Manner. G. G. Gould1, C. Smolik1,2, T. Gu1, M. Virela1, M. E. Valdez1, J. Hender2, A. Sellier2, A. Guillard2 and S. T. Schultz1. 1Physiology, UTHSCSA, San Antonio, TX; 2Pharmacology, UTHSCSA, San Antonio, TX; 3Family and Community Medicine, UTHSCSA, San Antonio, TX.

Pediatric acetaminophen use has increased since 1980, while baby aspirin use declined after its association with Reye Syndrome. It has been hypothesized that increased exposure of children under 5 years old to acetaminophen may be contributing to the rise in autism incidence. Cumulative acetaminophen doses < 75 mg/kg/day for 2 days are within recommended guidelines, but may be problematic for children in whom conjugative disposition pathways are impaired. Alternative metabolic pathways, either via deacetylation to promote endogenous cannabinoid neurotransmission or via oxidative pathways, may be harmful at vulnerable stages of brain development. We sought to model this scenario to clarify possible mechanisms using juvenile mice. We hypothesized that high-dose (>100 mg/kg/day) acetaminophen exposure in weanling (PD18-21) mice would produce impairments in the social behavior of C57BL/6j mice and would worsen them in BTRR mice. Male pups of each strain were exposed for 4 days to 4x daily injections of acetaminophen (100 mg/kg), WIN 55,212 (0.1 mg/kg) or vehicle (saline + 10% DMSO), and matured to adulthood (PD 80). Three chamber sociability tests of C57BL/6j mice revealed a significant (p<0.05) reduction in social interaction relative to vehicle controls that was comparable to prenatal treatment with valproic acid. However, the social behavior of BTRR mice was unaffected. Levels of 2-archidonoylglycerol and serotonin transporter binding site density were also reduced in both strains. In contrast C57BL/6j mice exposed for 4 days to 1.5 mg/ml of acetaminophen were unaffected. Thus, exposure to therapeutic doses of acetaminophen or cannabinoid full agonists indicates a possible role for cannabinoid receptors in modulating social behavior, at least in juvenile mice. This research was supported by US Navy contracts and a sub-award from UTHSCSA Institute for Integration of Medicine and Science CTSA grant #UL1RR025767.
Dogs demonstrate slower clearance and greater inhibition of DCA metabolism than rodents and most humans. The plasma kinetics of DCA in dogs is similar to humans with GSTZ1/MAI polymorphisms that confer exceptionally slow plasma clearance. Thus, dogs may be a useful model to further investigate the toxicokinetics of DCA.

Minipigs are often used in safety assessment studies with pharmaceuticals. Based on breeding experience, female minipigs are considered to reach sexual maturity at 4.5 months of age. However, more exact scientific data on sexual maturity is lacking. This information is pivotal for the design and interpretation of toxicity studies during drug development.

First, a pilot study was initiated to find useful parameters for the detection of estrous cycle in adult female Göttingen minipig. Three sows at 10-11 months of age, housed adjacent to adult boars were monitored daily for signs of heat (specific behavior, changes of the vulva) for 29 days. In addition, rectal body temperature was recorded, vaginal smears were taken for estrous cycle determination and the reproductive hormones progesterone and 17p-oestradiol were measured in serum. Progesterone analysis was the most valuable method, showing a cyclic pattern of release. The outcome of the examination of vulva and vaginal smear was less clear. Other external signs of heat did not reveal a cyclic pattern.

Subsequently, the approximate age at which female minipigs reach onset of sexual maturity was determined. For this, two gilts at 2 months of age, housed in the same room as adult boars, were used. Investigations included progesterone analysis, observations of the vulva, estrous cycle determination and recording of body weights. Progesterone levels increased first at approximately 5.5 and 6.5 months of age (body weight: 12.7 and 17.3 kg), respectively. The hormone release was indicative for a functional corpus luteum. For one female progesterone analysis was continued for nearly 3 months. The cycle duration was approximately 3 weeks which is in line with published data (18-21 days). However, the different stages of the cycle could not easily be distinguished by evaluation of vaginal smears. This information will be used in a following study to further investigate onset of sexual maturity in a female minipig population.

The purpose of this work was to evaluate a simple electroretinographic protocol on a representative sample of Göttingen minipigs. Electroretinogram recordings were conducted on 162 healthy minipigs (81 males and 81 females) aged 4–6 months. After a 1.5-hour light adaptation period, the animals were placed under general anesthesia. First, binocular full-field photopic electroretinogram recordings were conducted under photopic conditions. Subsequently, scotopic electroretinogram recordings were conducted during dark adaptation every 4 min over a 20-min period. At the end of this period, the maximal combined rod-cone response was recorded by measuring the retinal response to a single high-intensity flash. We used scleralcorneal clips as active electrodes and needle electrodes for the reference and ground. The a-wave and b-wave peak times and amplitudes were measured and statistically analyzed. For each of the statistical comparisons, normality and homogeneity of variances were evaluated. No significant gender differences were observed, with the exception of a higher b-wave amplitude for the photopic ERG recordings in females (48.14 ± 12.91 IV vs. 42.88 ± 10.67 IV; p < 0.005). The process of dark adaptation was evaluated, and the maximal combined rod-cone response was measured (a- and b-wave amplitudes and peak time). In conclusion, photopic and scotopic electroretinogram recordings were performed in the minipigs using a protocol based on light adaptation followed by dark adaptation. Scleralcorneal clip electrodes allowed a quick assembly and examination.

The miniaturization of the Hanford miniature swine, we report expanded and updated physiological data from normal intact and naive juvenile and young adult miniature swine of both genders. The normal physiological data gathered includes growth parameters, hematology, serum chemistry, coagulation profile, urinalysis, ECG rhythm and segment intervals, and organ weights.

Transgenic rodent gene mutation assays, recently adopted as OECD test guideline TG488, are useful methods to evaluate in vivo mutagenicity of chemicals in a target organ. The multiple copies of genetically neutral transgenes that contain reporter genes for mutation analysis are integrated in the chromosome of the transgenic rodents. However, biological identities between the transgenic and non-transgenic animals were not well demonstrated. To evaluate the characteristics of WistarHannover gpt delta transgenic (Tg) rat in general toxicity study, we investigated the clinical and pathological features of the gpt delta transgenic and non-transgenic WistarHannover rats in six months feeding study. The six-week-old Slc:WistarHannover/Rcc (gpt delta) and Slc:WistarHannover/Rcc (40 animals/sex/strain) were employed for this assessment. Twenty animals in each sex were subjected to necropsy at 13 and 26 weeks after starting, then clinical and pathological assessments were conducted. In Slc:WistarHannover/Rec (gpt delta), the values of several parameters in hematology, clinical biochemistry and organ weights were different from those in Slc:WistarHannover/Rcc with statistical significance. These alterations, however, were within the range of historical control data of in housed Slc:WistarHannover/Rcc, and thus were considered to be incidental. In histopathology, spontaneous lesions in WistarHannover such as fibrosis in the heart, microgranuloma in the liver and so on were observed in both strains. These results suggested that there were no remarkable differences which indicate phenotypic modulation in Tg rats, and supported the conclusion that Slc:WistarHannover/Rec (gpt delta) has similar characteristics to WistarHannover.
Comparison of Ophthalmologic Findings in Young Adult Crl: CD SD and Wistar Han Rats.


With more than one strain of rat now being used in toxicology studies, a study was instigated to see if there were differences in the incidence and nature of background lesions between the Crl: CD SD and the Wistar Han rat. The results of pretest ophthalmologic examinations of Wister Han (Charles River USA) and Crl:CD rats purchased from Charles River USA and BioLASCO Taiwan Co., LTD respectively were analyzed. The percentage incidences (total number of lesion (finding)/total number of rats) of the findings were compared for evidence of any significant differences between the two strains and sexes to see if any particular finding was unique or predominately expressed in one of the two strains/sexes. Corneal opacity along with the palpebral fissure is commonly observed in rats. Wister Han rats showed both a dense and higher incidence of this lesion when compared with Crl: CD rats. In addition, for Wister Han rats a higher incidence was observed for females as compared with males. This gender difference was not apparent for Crl: CD rats. Based on the differences of corneal opacity, the number eyes of Crl: CD rats that were considered normal was higher than that recorded for Wister Han rats. In addition, unilateral or bilateral posterior cortical multifocal pinpoint opacity was observed for Wister Han rats but not for Crl: CD rats. In conclusion, there is a clear difference in the incidence of spontaneous ocular findings observed for both Wistar Hans and Crl:CD rats; with Crl:CD rats showing a lower incidence of corneal opacity along with interpalpebral fissure and no posterior cortical lens opacity.

Feasibility of 3-Month Intravenous Bolus Injections via the Femoral Vein and an External Access Port in Sprague-Dawley Rats.

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Long term intravenous (IV) dosing continues to be critical in non-clinical development of many drugs (e.g., anti-cancer agents, large molecules), or pharmaceuticals that cannot achieve high systemic exposure by other routes. The IV route is commonly used in Sprague-dawley rats and the most common injection site is the tail vein. Although daily tail vein injection is normally feasible in acute and sub-acute studies (1, 7 or 14-day duration), sub-chronic and chronic studies (28-days, ≥3-months duration), once, twice and even multiple daily administration may be technically challenging and require additional animals at each dose level to assure an adequate number of animals to complete the study. Vehicle selection can also influence the success of long term daily or multiple daily tail vein injections and the use of the vena cava allows for a greater pool of blood to minimize secondary effects caused by non-ideal formulations. In alignment with the 3Rs (Reduction, replacement, refinement), our laboratory has demonstrated that daily intravenous injection via the femoral vein in the rat and external access port was feasible over a period of up to 3 months, which helped minimizing the number of animals per dose level to the minimum required by regulations. A medical grade polyurethane-based catheter was surgically implanted in the vena cava via the femoral vein and connected to a Quick-Connect harness with a Luer Valve. A heparin-based locking solution was used to fill the catheter and the injection cap. Sodium chloride for injection (0.9% Saline) was injected daily for up to 3 months via the injection cap, at a dose volume of 5 mL/kg. The experimental procedures were well-tolerated and no adverse effects were observed in clinical signs, body weight, and clinical pathology. Histopathological tissue changes were comparable to findings in catheterized animals. The IV bolus injection via the femoral vein and an external access port was shown to be a suitable alternative to tail vein injection for rat sub-chronic and chronic studies.

Noninvasive Airway Resistance Measurements in Large Animals by High-Frequency Airwave Oscillometry.

T. F. Schuessler1, A. Robichaud2, R. Forster3, M. Pouliot4, A. Ascalie, T. E. Tracey1 and S. Authier1, 1SCIREQ Scientific Respiratory Equipments, Montréal, QC, Canada; 2CiTeX LAB North America, Laval, QC, Canada; 3Faculty of Veterinary Medicine, University of Montréal, St-Hyacinthe, QC, Canada.

Large animals are generally more sensitive to rodents to drug induced respiratory changes. Methodologies to measure airway resistance in large animals normally require non-invasive techniques and compounds that can be orally administered or surgically implanted pleural pressure catheter. Both approaches could alter experimental endpoints in toxicity studies, which limit their use in drug development. A
non-invasive methodology to measure airway resistance by high frequency airwave oscillometry in anesthetized Beagle dogs and cynomolgus monkeys. A forced oscillation system was used to assess airway resistance and obtain conventional respiratory parameters (tidal volume, respiratory rate) before and after an intravenous treatment with a bronchoactive agent. Respiratory mechanics measurements were performed using a 16 second long signal (19 Hz) or composite (6, 10, 14, 19, 26 and 31 Hz) high frequency waveforms, which were applied at each time point at the subject’s airway opening via a face mask. During measurements, pressure and flow signals were recorded. After collection of baseline measurements, methacholine was administered to Beagle dogs (n=4) and cynomolgus monkeys (n=4) at 8 and 68 mcg/kg, respectively. A peak increase in airway resistance (+64% in Beagle dogs and +30% in cynomolgus monkeys) was observed after intravenous methacholine administration in both species with return to baseline comparable levels within 10 min. Airway resistance data analysis for individual animals revealed mean percent variations from baseline ranging from 9.5 to 18.6% for dogs and from 4.4% to 6.9% for cynomolgus monkeys, suggesting stable and reproducible assessment of lung function. Airwave oscillometry appears to be suitable for non-invasive respiratory mechanics measurements in large animal toxicology studies.

Seizure liability studies in non human primates generally aim to: 1) confirm drug-induced seizures are self-limiting, 2) determine plasma level at seizure onset 3) identify prodromal clinical signs which can be monitored in clinical trials 4) confirm that conventional drugs (e.g. diazepam) can treat drug-induced seizure and 5) confirm the no observed adverse effect level (NOAEL) by absence of paroxysmal activity. To achieve these goals, typical study designs include video-EEG monitoring. Continuous telemetry monitoring generates a considerable amount of data which translates into lengthy analysis when solely relying on manual EEG review. A new application for seizure detection was developed in non-human primates (NHP). The prototype evaluates temporal and spectral features and compares them to features extracted from a sliding reference window. Detected seizures are validated based on a decision tree and artifact rejection algorithms. The automated prototype was used to evaluate periods of up to 68h of continuous EEG data and all detections were subject to a visual review. All animals presented at least one epileptic event. A detection sensitivity of 86.4% with 2.3 false positives per hour of signal was achieved using the default parameters of the prototype. Detection sensitivity increases to 100% when selecting optimal settings for each individual. The processing time was approximately 5 minutes for a 24h EEG telemetry signal. These promising results reflect progress in an era computerized data analysis. Further improvements are investigated to maintain a detection sensitivity of 100% essential with the model, while keeping the number of false detections low to increase data analysis efficiency.

Comparing Data Obtained by Refined Restraint Methods to Traditional Methods in Nonhuman Primate Studies.

As nonhuman primates continue to be an important model in nonclinical toxicology/pharmacology studies, improving methods of animal handling during procedures is essential. The aim of this study is to present historical background data collected using a refinement in the restraint technique, the Procedure Cage (PC). Use of the PC has the potential to contribute to minimizing inter- and intra-animal data variability when compared to data collected using other restraint methods (e.g. restraint chair [RC]). Historical control pregnancy loss ratio of 16.3% (range 0–31.3%) obtained using the PC is relatively low when compared to published data (17.8%, Hendrie et al. 1996, Am J Primat 40:41; and 29.6%, Wehner et al. 2009, BDR [B] 86:144). Restraint-associated stress can also influence values of hemodynamic and respiratory measurements. In a telemetry study comparing cardiovascular parameters for restraint in the Restraint Chair (RC) vs. Procedure Cage (PC), six telemetry implanted animals were monitored and data analyzed for heart rate (HR) and mean arterial blood pressure (BP) using a validated Notocord Telemetry System. In a series of procedures performed, HR for RC animals was on average 1.2 times (range 147–198 bpm) of BP for RC animals and was on average 1.2 times (range 139–167 bpm). BP measured from PC animals was on average 1.2 times (range 102–109) that of PC animals (range 90–96). The RC animals returned to baseline HR and BP levels only 22 and 33% of the time, beginning at 15 and 30 min post procedure, respectively. In contrast, the PC animals returned to baseline HR and BP levels 50 and 91% of the time beginning at 1 and 10 min post procedure, respectively. In vitro it is evident that use of the PC method has potential to contribute to obtaining more robust data and thus, could eventually lead to use of fewer animals on study.

Comparative Analysis of Noninvasive Blood Pressure Data by High-Definition Oscillometry (HDO) in Cynomolgus Monkeys of Mauritian and Asian Origin.

Cardiovascular investigations - among other diagnostic parameters – are essential for the selection and interpretation of untoward findings in preclinical studies. As cardiovascular toxicity in general and blood pressure determination in particular is concerned with adverse effects of xenobiotics on the circulatory system, a representative analysis of non-invasive blood pressure data is mandatory for the right interpretation of apparent circulatory findings. Cynomolgus monkeys are an attractive model in toxicity studies and animals from various sources such as Mauritius and Asia are available from acknowledged breeders. The aim of the present work was to compare non-invasive blood pressure data from group housed, untreated and unseated cynomolgus monkeys of Mauritian and Asian origin. Individual data from 150 Mauritian and 92/128 Asian male and female animals, each were analysed for systolic, diastolic and mean arterial blood pressure. Blood pressure data of male and female animals of either origin showed slight but significant differences (P ≤ 0.05 to P ≤ 0.0001) and values for males were higher than for female animals. Comparison of blood pressure data from monkeys of Mauritian and Asian origin revealed that systolic pressure (+9.0 % M; +10.4% F), diastolic pressure (+12.0 % M; +14.1 % F) and mean arterial pressure (+10.1 % M; +11.7 % F) were significantly (P ≤ 0.01 and P ≤ 0.001) elevated in monkeys from Mauritius when compared to monkeys from Asia.

Procedure Refinement and Reduced Restraint Enables Extended 6-Hour-Daily Inhalation Dosing in Beagle Dogs.

Extended duration of dosing for preclinical assessment of inhaled pharmaceuticals is considered advantageous when testing effects of expected low toxicity. Technical limitations have often determined the maximum time of exposure in non-rodents such as Beagle dogs. Increased exposure time can improve the chances and accuracy of determining the maximum tolerated dose rather than being limited to a maximum feasible dose. A study was undertaken to confirm whether reduced restraint (platform) combined with procedural refinements would enable routine 6 hr daily dosing (5 days) and even 6 hr daily dosing (3 days). Prior to delivery from the supplier, a number of dogs were presented to platform restraint and inhalation facemask for up to 15 min and those demonstrating acceptance were selected. Following receipt, the four dogs were gradually acclimated for increasing periods of time (up to 4 hrs) to the inhalation equipment over 13 consecutive days. During the acclimation period, the dogs were monitored for behavior changes (excessive salivation, trembling, vocalization, struggling and increases in respiratory rate). Reinforcement through positive behavior by voice and minimal animal contact during the sham dosing where emphasized. Verbal rewards, petting and treats were provided after completion of each session. All animals underwent the 4 hr daily sham dosing except one female for which the sham dosing had to be interrupted on Days 2 to 4. The 6 hr sham dosing was initiated and successfully conducted using the remaining 3 dogs. There were no changes in clinical signs, body weights or food consumption throughout the study. In conclusion, extended daily inhalation dosing of Beagle dogs is made possible by combining animal screening at the supplier, extensive acclimation, positive behavior re-enforcement and reduced restraint. This enables to more accurately determine the NOEL and improve the determination of a MTD while improving animal welfare by reducing the level of restraint and stress.
volume for hematology and clinical chemistry, we investigated dilution of small blood volumes collected by jugular venipuncture. Whole blood (300 μL) collected at 1:2 mL collected from 46 mice was split; 150 μL for hematology and 150 μL for clinical chemistry. For hematology, 2 blood and 2 reticulocyte smears were prepared prior to diluting 100 μL of blood with 200 μL of ADVIA sheath rinse solution®. The 1:3 dilution was analyzed on the ADVIA 120 hematology analyzer (Siemens). All flagged values were verified by microscopic blood smear evaluation. Serial blood dilutions yielded adequate precision up to 1:5 dilution. For clinical chemistry, 40 to 50 μL of serum was diluted 1:3 in saline. Samples were analyzed on the Modular Analytics biochemistry analyzer (Roche) with exception of electrolytes. Serial dilutions of all chemistry parameters evaluated had previously demonstrated acceptable precision up to 1:5 dilution in saline. Jugular vein versus abdominal aorta sample results, were compared using historical ranges. Twelve of 46 (hematology) and 8 of 46 (clinical chemistry) samples were inadequate for analysis, similar incidence to abdominal aorta sampling. Jugular vein sampling had slightly higher RBC and platelet counts, while clinical chemistry parameters were comparable except for slight elevations in aspartate aminotransferase. These dilutions necessitated more technical sample processing, but allowed clinical pathology analysis with only 300 μL of blood (ie. 4X reduction), presenting clear advantages, including interim study sampling, reduction in animals, combination of endpoints for in a given mouse, improving correlations and interpretation.

**S 765 An Assessment of Exposure Pathways and Potential Impacts to Human Health Associated with Shale Gas Drilling and Production Operations.**

A. Harris. ENVIRON International Corporation, Little Rock, AR.

Shale gas drilling operations continue to expand in the US, leading to increasing focus on identifying and understanding potential impacts to human health. The primary concern is the potential contamination of air and groundwater in communities located near shale gas drilling sites. Air quality may be affected from drilling and fracturing activities as well as emissions from vehicles and equipment. Air quality studies have been conducted both at drilling sites and near community receptors. VOCs, when detected, have been at concentrations below health-based benchmarks. Multiple studies have also been conducted to evaluate potential impacts to groundwater by additives of fracturing fluid, petroleum constituents and methane. Migration of fracture fluids to groundwater aquifers have been proposed although there is currently little data indicating that groundwater has been impacted by fracture fluids. Other exposure pathways of concern include leaks or spills of undiluted additives to surface water and inadequate containment of flowback and produced wastewater.

**S 766 The Potential Toxicological Impacts of Shale Gas Drilling: An Overview.**

B. Goldestein and J. Kreisky. University of Pittsburgh, Graduate School of Public Health, Pittsburgh, PA.

The pace of development of hydraulic fracturing technology to extract natural gas from shale formations has outstripped that of the research necessary to ensure that we maximize the benefits and minimize the risk of this important activity. Issues of particular toxicological concern include the potential impact of the evolving list of chemical and physical agents which are used in hydraulic fracturing. Concerns also extend to upstream processes, such as the mining of silica and the delivery of chemical and physical agents to the site; the on-site drilling, hydraulic fracturing and long-term extraction processes, including contamination of groundwater, and emissions and noise from diesel compressors; and downstream activities such as the distribution of the hydrocarbon products and the safe disposal of the literally millions of gallons of fluid that comes up from underground containing such agents as brine components, arsenic and radioactive agents. Of particular toxicological importance is the potential adverse consequences of the mixtures of hydrofracturing agents, hydrocarbons and natural agents brought to the surface. Air pollution also has been documented and there is concern about the nearby impacts of the release of hydrofracturing agents and volatile hydrocarbons, as well as regional ozone issues. The longer term legacy of site specific pollution is also an issue.

**S 767 Community Exposure and Risk Near Natural Gas Production Sites: Impacts and Research Needs.**


The current boom in domestic natural gas production has focused national attention on the environmental and health impacts of this process. Increased development of unconventional natural gas resources (i.e., from shales, coal beds and tight sands) using hydraulic fracturing (“fracking”) has raised concerns about impacts on environmental quality and human health, especially as development has moved into populated areas. While public concerns tend to focus on potential water contamination, the well development process also results in emissions of multiple air toxics when injected water “flows back” to the surface with the natural gas, accompanying petroleum condensate and fracturing chemicals. Studies in Colorado and elsewhere indicate that multiple air toxics and chemicals used in well production activities are emitted from gas development and production sites. Research on non-occupational exposure indicates that air pollution is a major concern because health-based risk estimates for nearby residents are greater than for individuals living further from well development sites. Human exposures and risks vary with proximity to wells, specific activities at well pads, topography/meteorology, and individual operator practices and controls. This talk explores our current understanding of exposure and health risks around well development sites, identifies major information gaps and uncertainties, and outlines research that will improve our understanding of the health risks associated with natural gas development.
Texas is home to the Barnett Shale, a shale-containing geological formation in a highly-populated area which is currently being actively drilled using hydraulic fracturing. The Texas Commission on Environmental Quality (TCEQ) is charged with regulating sources of air emissions from natural gas operations and has spent considerable resources examining emissions from these sources. The TCEQ has utilized flyovers of helicopters outfitted with infra-red (IR) cameras capable of visualizing volatile organic compound (VOC) emissions, conducted numerous mobile monitoring trips, and has installed an extensive ambient air fixed-site monitoring system in areas with intensive drilling. Between August 1, 2009, and June 30, 2012, the TCEQ has surveyed 2,307 sites in the Barnett Shale area using hand-held IR cameras, and at 2,263 of these sites, hand-held survey instruments were also used. These surveys included citizen complaint investigations, compliance investigations, mobile monitoring trips, and follow-up investigations based on helicopter flyovers. As a result of observations with the hand-held IR camera and measurements collected using the survey instruments, 1,167 short-term field canister samples have been collected and analyzed for VOCs. Less than 5% of VOC canister samples measured short-term levels of concern. Short-term samples have also been collected and analyzed for carbonyls, NOx, and sulfur compounds and none of the analyses have measured chemicals at short-term levels of concern. Six stationary VOC monitors were operational in the Barnett Shale area in 2009, thirteen are operational now. No VOCs have been detected to date at levels of long-term health concern.

Considerable resources have also been devoted to developing mobile monitoring systems that can detect air quality impacts of natural gas operations in Texas. In early 2010, the TCEQ commissioned an additional mobile monitoring system with on-board long-term canister samplers to collect VOC samples that could be analyzed for long-term effects. This mobile monitoring system is equipped with a hand-held IR camera, long-term canisters, and a fixed-site air quality monitoring station. These canisters are placed in the areas of interest and then collected and analyzed for VOCs. This system is being used to study the long-term impacts of natural gas operations on air quality in areas with intensive drilling. Between August 1, 2009, and June 30, 2012, the TCEQ has deployed this mobile system at 1297 sites in the Barnett Shale area. At 1290 of these sites, hand-held survey instruments were also used. These surveys included citizen complaint investigations, compliance investigations, mobile monitoring trips, and follow-up investigations based on helicopter flyovers.

As a result of observations with the hand-held IR camera and measurements collected using the survey instruments, 1,167 short-term field canister samples have been collected and analyzed for VOCs. Less than 5% of VOC canister samples measured short-term levels of concern. Short-term samples have also been collected and analyzed for carbonyls, NOx, and sulfur compounds and none of the analyses have measured chemicals at short-term levels of concern. Six stationary VOC monitors were operational in the Barnett Shale area in 2009, thirteen are operational now. No VOCs have been detected to date at levels of long-term health concern.

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**Role of Metabolic Syndrome and Perivascular Adipose in Exposure-Induced Vascular Dysfunction.**

D. J. Conklin and M. J. Campeu.

Recent epidemiological studies indicate that ambient air pollution enhances the progression from metabolic syndrome to diabetes but not by increasing obesity. Because vascular dysfunction and vascular insulin resistance are two early events associated with metabolic syndrome and air pollution exposure, the mechanisms by which these alterations occur in the vasculature could be informative of changes in overall cardiovascular disease risk. Similarly and increasingly, there is a growing recognition that the perivascular adipose tissue (PVAT) is an important regulator of vascular tone under normal homeostatic conditions and PVAT is altered during disease states, including metabolic syndrome and diabetes. The perivascular adipose tissue structure and function are changed to a proinflammatory state both by disease, angiotensin II, and by exposure to environmental pollutants, implicating these alterations in subsequent endothelial and vascular dysfunction. This symposium will provide both an introduction to the role of perivascular adipose tissue in health and metabolic disease states and draw on evidence from environmental and experimental model studies to emphasize specific alterations in perivascular adipose tissue and how these changes contribute to subsequent vascular dysfunction and potentially increase the risk of cardiovascular disease.

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**Turning Vascular Disease Inside Out: Role of Perivascular Adipose Tissue in Coronary Artery Disease.**

N. Weintraub. Division of Cardiometabolism, University of Cincinnati, Cincinnati, OH. Sponsor: D. Conklin.

Adipose tissue surrounding the great vessels [perivascular (PV) adipose tissue, or PVAT] expands during obesity and correlates with the extent of visceral fat and insulin resistance in humans. Moreover, both the amount of PVAT and the degree of PVAT inflammation correlate with the presence of atherosclerotic coronary artery disease. Studies in isolated perivascular adipocytes indicate that the cells display a reduced state of adipogenic differentiation, and enhanced pro-inflammatory cytokine expression, as compared with subcutaneous and visceral adipocytes. In response to just two weeks of high fat feeding in mice, adipogenic gene expression is markedly reduced, while pro-inflammatory cytokine expression is up-regulated, in PVAT. Using a novel model of PVAT transplantation to the mouse carotid artery in the setting of high fat feeding, we demonstrate that PVAT enhances atherosclerosis and wire injury-induced neointimal hyperplasia, triggering inflammatory cell infiltration and proliferation of vasa vasorum. Our data suggest that PVAT plays a pathogenic role in vascular disease, linking metabolic signals to inflammation and angiogenesis of the blood vessel wall.

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**Perivascular Adipose and Vascular Dysfunction in Mice after Combined High-Fat Diet and Concentrated Ambient Particulate Matter Exposure.**

D. J. Conklin and P. Haberzettl, Cardiovascular Medicine, University of Louisville, Louisville, KY.

Exposure to inhaled fine particulate matter induces endothelial dysfunction and increases the risk of cardiovascular disease, insulin resistance and the risk of diabetes, however, the mechanisms by which PM2.5 enhances this risk are unclear. Because diabetes and air pollution are worldwide health problems, we examined whether high fat diet (HFD) enhanced concentrated ambient particulate matter (CAP)-induced EPC suppression and insulin resistance to ascertain the relationship between these important changes. Mice (male; C57BL/6) maintained on low fat (LFD, 10% kcal fat) or high fat (HFD, 60% kcal fat) were exposed to HEPA-filtered air or urban Louisville CAP (80-100 μg/m3) for 9 or 30 consecutive days (6/6d). CAP exposure or HFD prevented insulin-induced Akt and eNOS phosphorylation in isolated aorta, and aortic contractility was progressively dysfunctional under CAP, HFD and combined HFD+CAP treatments. Moreover, only combined HFD+CAP treatment increased glucose intolerance without increasing obesity, and thus, worsened the metabolic syndrome and progression toward diabetes. These data implicate alterations in vascular structure such as increased peri-vascular adipose in the combined effects of HFD+CAP on vascular dysfunction and insulin resistance.

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**NADPH Oxidase and Perivascular Superoxide Production with Particulate Matter Exposure.**

Q. Sun. Environmental Health Sciences, The Ohio State University, Columbus, OH.

Reactive oxygen species from oxidant-releasing enzymes such as NAD(P)H oxidase are known to play a causal role in certain disease development, and air pollutants, such as PM2.5, are known to cause vascular dysfunction in major cardiovascular diseases. To evaluate the role of exposure to airborne fine particulate matter (diameter, <2.5 μm, PM2.5) pollution on metabolic parameters, inflammation, and adiposity, PM2.5 inhalation exposure was performed in C57BL/6 mice and mice deficient in the cytosolic subunit of the nicotinamide adenine dinucleotide phosphate (NADPH) oxidase p47phox (p47phox−/−) with normal or high fat diet. We found that PM2.5-exposed C57BL/6 mice exhibited metabolic abnormalities consistent with insulin resistance. Ex vivo-labeled and infused monocytes demonstrated increased adherence in the microcirculation of normal diet- or high-fat diet–fed PM2.5-exposed mice. p47phox−/− mice exhibited an improvement in parameters of insulin resistance, vascular function, and visceral inflammation in response to PM2.5. We concluded that exposure to high levels of PM2.5 is a risk factor for subsequent development of insulin resistance, adiposity, and inflammation, and reactive oxygen species generation by NADPH oxidase appears to mediate this risk. Other NAD(P)H oxidase substrates and their roles in PM2.5 exposure-induced cardiovascular diseases will also be summarized.

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**Role of Perivascular Adipose in Regulation of Vascular Function by the Aryl Hydrocarbon Receptor.**

M. K. Walker. Department of Pharmaceutical Sciences, University of New Mexico, Albuquerque, NM.

The aryl hydrocarbon receptor (AHR) has been demonstrated to be required for normal blood pressure regulation as demonstrated by the hypotensive phenotype in the global AHR knockout mouse. Notably, conditional deletion of the AHR solely from endothelial cells results in a nearly identical blood pressure phenotype as observed in the global knockout. Thus, we hypothesized that AHR plays a critical role in the vascular regulation of blood pressure. We generated ECahr−/− mice by crossing AHR Rosed mice (ahrfl/fl) to mice expressing Cre recombinase driven by an EC-specific promoter. BP was assessed by radiotelemetry prior to and following an acute injection of the potent vasocostricton, angiotensin (Ang) II, or chronic treatment with an inhibitor of Ang II formation, Ang converting enzyme inhibitor (ACEi). ECahr−/− mice exhibited significantly different responses to Ang II and ACEi. While Ang II increased BP in both genotypes, the increase was sustained in ECahr+/+, whereas the increase in ECahr−/− mice steadily declined. Area under the curve (AUC) analysis showed that Ang II-induced increase in diastolic BP (DBP) over 30 min was significantly lower in ECahr−/− mice (AUC: ECahr+/+ 129±223 mmHg/30 min; ECahr−/− 50±138 mmHg/30 min, p<0.05). In contrast, while ACEi decreased BP in both genotypes, the subsequent rise in DBP after treatment was significantly delayed in the ECahr+/+ mice. ECahr−/− mice also exhibited reduced vascular and adipose Ang II type 1 receptor (AT1R) expression, and reduced aortic Ang II-dependent vasocostriction in the presence of vascular
Adipocytes express all components of the renin-angiotensin system necessary to synthesize and respond to angiotensin II (AngII). We have previously demonstrated that infusion of AngII to hyperlipidemic male mice augments atherosclerosis and causes the formation of abdominal aortic aneurysms (AAAs). Mechanisms for AngII-mediated AAAs include inflammation that extends to perivascular adipose tissue surrounding the abdominal aorta. Inflammation in perivascular adipose tissue was augmented by genetic or diet-induced obesity, and resulted in marked increases in susceptibility to AngII-mediated AAAs. Moreover, weight loss in previously obese mice resulted in remodeling of AAAs. Vascular changes in mice experiencing weight loss included reduced adventitial neovascularization, with alterations in stem cell populations in perivascular adipose tissue. Current studies are examining effects of deficiency of components of the renin-angiotensin system in adipocytes on AngII-mediated vascular diseases.

Adipocytes are implicated in the etiology of protein conformational diseases, including Parkinson’s disease (PD) and Alzheimer’s disease (AD). The purpose of this workshop is to bring together experts to discuss etiologic factors associated with PD and AD, particularly environmental pesticides, heavy metals, and other pollutants, and to discuss new directions for identifying suitable targets for therapeutic strategies. Two very promising lines of investigation into the mechanisms of dopaminergic cell death in PD will be discussed, as well as the interaction of senescence-related, genetic, and environmental factors likely contributes to the etiology of late-onset neurodegenerative diseases such as PD and AD. The interactions of heavy metals with chaperone proteins that regulate protein folding and cell death as related to protein conformational diseases will also be discussed. The workshop will address six questions: 1) Despite equivocal results, why are genetic factors still considered the primary etiological factor in late-onset neurodegenerative disease?; 2) How can existing models of neuronal cell death in neurodegenerative diseases be integrated?; 3) How can cell culture models be integrated into a better understanding of neurodegenerative disease processes?; 4) Are there common pathways or mechanisms that are relevant to environmental exposures in PD and other neurodegenerative diseases?; 5) What is the evidence that early exposures to environmental factors predispose to neurodegenerative diseases?; 6) What is the evidence that environmental factors associated with neurodegenerative diseases? (supported by NIH grants ES10586, NS 074443, ES19267).
pathology by disrupting the normal storage and release of dopamine and related neurotransmitters. Dr. Miller will also present novel high throughput approaches that will be used to further study this association. Supported by NIEHS P01ES016731.

### W 779 Integrating Epidemiological and Basic Research to Assess the Role of Pesticide Exposure in Neurodegenerative Disease.

**L. Richardson, Robert Wood Johnson Medical School, Piscataway, NJ.**

Although a significant amount of research effort and resources have been focused on determining genetic contributors to Parkinson's disease (PD) and Alzheimer's disease (AD), purely genetic contributions account for only a fraction of the cases. Our laboratory has been specifically interested in the potential role of pesticides in the etiology and pathogenesis of both PD and AD. Recently, we have found that serum levels of a long-lasting residue of the organochlorine pesticide hexachlorocyclohexane (B-HCH) was significantly higher in the serum of PD patients, while the DDT metabolite, p,p'DDE, was significantly higher in patients with AD. DDE levels were 3.8-fold higher in serum of AD patients (2.61 ± 0.35 ng/mg cholesterol; n = 86) when compared to controls (0.69 ± 0.10 ng/mg cholesterol; n = 79; p < 0.001). After controlling for age, sex, education, race, and sample collection site, the OR for increased risk of AD in the highest tertile of DDE levels was 4.26 (95% CI: 3.41-5.32; p < 0.001). To determine whether DDT exposure alters the expression of AD-related genes in vivo, male C57Bl6 mice were dosed with 3 mg/kg DDT every 3 days for 30 days. DDT significantly increased the mRNA levels of APP (25%), NEP (77%), and ApoE (150%) in the hippocampus. These data suggest that DDT selectively alters the expression of AD-related genes in a brain-region specific manner with the hippocampus being more sensitive than the frontal cortex. Specific Questions to be addressed will include: Despite equivocal results, why are genetic factors still considered the primary etiological factor in late-onset neurodegenerative disease? Are there common pathways or mechanisms that are relevant to environmental exposures in both PD and AD? How do chemicals in a similar class of compounds elicit very different effects on the neurodegenerative processes?

### W 780 Life-Course Models for Ensuring Children's Health Protection.

**S. P. Darney** and E. M. Faustman. 1 ORD, US EPA, Research Triangle Park, NC; 2Environmental & Occupational Health Sciences, University of Washington, Seattle, WA.

New knowledge about environmental risks to human reproduction and development directly relevant to children’s health protection derives from the fields of developmental and reproductive toxicology, exposure science, epidemiology, risk assessment, and public health. Together, this information highlights the importance of the intrauterine environment in setting the stage for lifelong health, along with the complexities of the physical, chemical, and social factors that operate during critical windows of development to impact health and wellbeing. For example, breakthroughs in genetic polymorphisms and epigenetics are extending our understanding of inherent and acquired susceptibility to effects of environmental contaminants and showing how various intrauterine stressors such as nutrition, toxicants, and social stress may alter developmental programming at the start and throughout life. These scientific advances point to the need for innovative cumulative risk assessment methods and public health intervention approaches in order to account for risks that accrue across the developmental continuum from cradle to cradle. This workshop brings together the interdisciplinary expertise needed to begin integrating new knowledge into life-course models for children's health and wellbeing. Topics include research findings from toxicity testing and epidemiology studies specific to critical windows of exposure during pre-conception, pregnancy, and early childhood and strategies to stimulate research on and optimizing testing and risk assessment models and enabling analyses across the whole life course. The workshop also features an innovative approach for evaluating and communicating complex scientific information about reproductive risks and interventions to diverse audiences including health care providers, parents (present and future), and regulators.

### W 781 Today's Challenge for Protecting Children's Environmental Health for a Lifetime.

**E. M. Faustman. University of Washington, Seattle, WA.**

Evidence continues to build for the increasing importance of early life experiences in defining not only childhood development and normal health trajectories but also the potential for chronic disease risk throughout later life. Thus, a critical need for protecting children's health is recognition that our models require a dynamic context--a life course framework in order to address the key differences in development of environmental contaminants and showing how various intrauterine stressors such as nutrition, toxicants, and social stress may alter developmental programming at the start and throughout life. These scientific advances point to the need for innovative cumulative risk assessment methods and public health intervention approaches in order to account for risks that accrue across the developmental continuum from cradle to cradle. This workshop brings together the interdisciplinary expertise needed to begin integrating new knowledge into life-course models for children's health and wellbeing. Topics include research findings from toxicity testing and epidemiology studies specific to critical windows of exposure during pre-conception, pregnancy, and early childhood and strategies to stimulate research on and optimizing testing and risk assessment models and enabling analyses across the whole life course. The workshop also features an innovative approach for evaluating and communicating complex scientific information about reproductive risks and interventions to diverse audiences including health care providers, parents (present and future), and regulators.

### W 782 Periconception Parental Metal Exposures, Couple Fecundity, and Child Health: Delineating the Chicken and Egg Question.

**G. M. Buck Louis, R. Sundaram and J. Maisog. Division of Epidemiology, NICHD, Bethesda, MD. Sponsor: S. Darney.**

Growing evidence suggests that human fecundity and fertility may be informative about the subsequent health and well-being of couples’ offspring. This avenue of study is in keeping with the early origin for health and disease hypothesis (DOHaD), and the need for a life course approach for assessing health across the lifespan. Early evidence suggests that gynecologic and urologic disorders may be associated with adverse pregnancy outcomes such as genital-urinary malformations or diabetes in affected children. Similarly, couples experiencing conception delays are reported to be at higher risk of delivering infants of diminished gestation and birth size, outcomes associated with developmental disabilities and other childhood morbidities. This evolving body of evidence underscores the importance of exposure assessments in both partners in keeping with the couple-dependent nature of human reproduction and development. For illustrative purposes, this talk utilizes data from the recently completed Longitudinal Investigation of Fertility and the Environment (LIFE) Study to demonstrate the relation between couples’ exposures to persistent environmental chemicals (i.e., cadmium, lead, and mercury) and fecundity as measured by time-to-pregnancy (in menstrual cycles), and infant health outcomes such as gestation and birth weight. Blood cadmium concentrations in female partners (range 0.02-2.87 μg/L) and blood lead in male partners (range 0.34, 6.91 μg/dL) were significantly associated with a longer time to pregnancy as measured by fecundity odds ratio (FORS) below one (FORS<0.78; 95% CI 0.63, 0.97 and FORS<0.85, 95% CI 0.73, 0.98, respectively). When jointly modeling both partners’ metal exposures as a continuous exposure, female lead and male cadmium exposures were associated with reductions in gestation and birth weight. Further analysis is underway to determine if the reduction in gestation and birth weight is mediated indirectly through reduced couple fecundity or directly on fetal growth and development.

### W 783 Prenatal Pesticide Exposures and Children's Neurodevelopment across the Life-Course Continuum—Lessons from the Fields.

**B. Eskenazi, A. Bradman, K. Harley and N. Holland. Center for Environmental Research and Children's Health, University of California Berkeley, Berkeley, CA. Sponsor: S. Darney.**

A growing body of evidence suggests that prenatal pesticide exposure may be related to adverse effects on neurodevelopment of children. In 2000-2001, we initiated CHAMACOS (Center for the Health Assessment of Mothers and Children of Salinas), a longitudinal birth cohort study, with the primary aim to investigate the association of prenatal and postnatal exposure to agricultural chemicals and health and development of children. We enrolled 601 pregnant women into this community-based participatory research project. The participants were primarily low-income Latinas from the agricultural community in the Salinas Valley, Monterey,
California. We conducted detailed neurodevelopmental assessment of the CHAMACOS children at multiple time points and currently the children are reaching their 12th birthday. During these years, we have monitored their exposures to a number of chemicals including pesticides such as organophosphates, organochlorines and fungicides. We have also examined potential genetic susceptibility to exposure with analyses of the PON1 polymorphisms and enzyme activity and we are now beginning analyses of epigenetic endpoints. These studies highlight the advantages of longitudinal cohort studies for evaluating the long-term impact of prenatal and early life exposures on children’s health and development. Results of the CHAMACOS study to date, coupled with those of other longitudinal studies demonstrate how in utero exposure to certain pesticides can impact neurodevelopment.

Furthermore, our results suggest that certain children by virtue of their genetic make-up maybe more susceptible. Lastly, we have identified limitations in assessing exposures to pesticides that challenge our ability to determine “causal” relationships. Primary funders: Children’s Center grant from NIEHS, NIH and US EPA.

### New Strategies for Addressing Toxicity Testing across the Lifespan

**P. M. Foster, National Toxicology Program (NTP), NIEHS, Research Triangle Park, NC**

A previous NTP workshop evaluating the utility of the cancer bioassay for detecting hormonally-related cancers (particularly breast, prostate, ovary and testis), concluded that the standard approach, commencing exposure as young as adults (6-9 weeks of age) is likely missing important windows during early development. It could be critical for cancer outcomes and therefore assessment of lifetime risk. NTP used the workshop results to adopt a new default for rat cancer bioassays to incorporate early life exposures into the assay. NTP had previously conducted perinatal bioassays, but required specific justification; now, such studies are conducted routinely, unless there is a scientific rationale not to do so. This study begins exposure at implantation and continues until the offspring reach 2 years of age. Embarking on a cancer study requires preliminary dose setting information. For a cancer bioassay, this is normally a 90 d study, but would now require exposure from implantation thru weaning prior to the 90d component to evaluate target organ toxicity. Such an approach, would also lend itself easily to evaluations of multiple other endpoints from the same exposure paradigm, including developmental neurotoxicity and immunotoxicity, and, if one bred the F1 offspring at adulthood, both fertility/ fecundity and pre-natal developmental toxicity (teratology) - the NTP modified one-generation study. Each of these endpoints could be considered interchangeable “cassettes” depending on the type of toxicity that NTP was required to assess. Compared to the individual, stand-alone studies, this would have considerable savings in time, cost and animal usage, yet generate high quality developmental and reproductive toxicity information for risk assessment. This is a similar approach to the OECD extended generation study, but has several major advantages, most importantly is a significantly improved ability to detect potential effects on fertility and fecundity in the same animals on which structural changes are evaluated, together with a pre-breeding exposure that is consistent with the length of rat spermatogenesis.

### Advancing Risk Communication and Decision Tools for Children’s Health Protection

**T. J. Woodruff and P. Sutton, Program on Reproductive Health and the Environment, UCSC, San Francisco, CA. Sponsor: S. Darnay**

Parents and physicians who take care of them and their children read and hear a plethora of information, some contradictory, about environmental and chemical risks to children’s health. Better tools are needed to evaluate and synthesize this information and translate it into meaningful prevention strategies for health professionals, the public and policy-makers. Methods to synthesize the science into evidence-based decision tools vary according to context, including individual, community, and population settings. They may be hierarchical or evidence-based decision tools. The potential for synthesis is enhanced through the identification of underlying quantitative relationships between their structural, chemical, and physical properties and responses in key biological systems. The Toxicity in the 21st century testing paradigm holds particular promise for nanoparticles because whole animal testing is not economically practical or feasible for every new material introduced to the marketplace. Important advances have been made in this emerging but critical field, yet many challenges remain. Distillation of the large amount of phenotypic data regarding in vitro cytotoxicity and inflammation down to in vivo-relevant adverse events is a daunting task, given the vast differences in microenvironment, temporal dynamics, and exposure. The use of systems biology approaches will enable the identification of relevant target organs/cell types, the development of mechanism-based assays for hazard assessment and computational modeling for predicting potential toxicity from human relevant exposures. This session will address computational challenges in data standards, data integration, and modeling of dose response and structure-activity relationships, including talks by experts in the areas of algorithms, in vitro screening, in vivo exposures and PB/PK modeling. Included is an opening by the co-chair of the US-EU Nanoinformatics Community of Research, which seeks to facilitate intercommunication between researchers, regulators, and grant agencies on environmental health and safety issues for manufactured nanomaterials. Speakers will include current experimental and computational approaches as well as future research needs to identify relevant in vitro assays that are predictive for nanomaterial hazard assessment based on in vivo mode of action and realistic human exposures.

### US-EU Nanotechnology Databases and Ontology Community of Research

**N. A. Baker¹ and H. Rauscher². ¹Pacific Northwest National Laboratory, Richland, WA; ²DG Joint Research Centre Unit Nanobioscience/TP 202, European Commission, Ispra, Italy. Sponsor: K. Waters**

This talk will provide an overview on the work of a new US-EU Nanotechnology Safety Community of Research (CoR) focused on nanotechnology databases and ontology with specific examples and highlights from our own research. CoRs are groups of people who share a significant interest in nano safety, work together to develop a shared repertoire of resources, and have regular contact to address key global research challenges. Interconnected information systems are urgently needed for collating nanoscale material descriptions; their intrinsic and context-dependent properties and their interactions; and their interactions with biological entities. One goal of the CoR is to enable the sharing, searching, and analysis of nanoscale material characterization data across a wide range of experimental sources and guide the structuring of these data to enable their widest possible use. Achievement of this goal will deliver important new capabilities to allow integration of risk assessment data among labs throughout the world, to provide situational awareness of data coverage across nanomaterial categories, and to enable predictive computational models for bridging physical properties and biological outcomes with exposure, dispersal and fate. In order to realize this goal, the CoR will initially focus on the following three areas of investigation: Identification of the data elements necessary to establish common data-sharing models for this domain; specification of requirements for sharing data between research groups and repositories in human- and machine-interpretable forms; definition of concepts necessary to support the above activities and representation of those concepts in an ontological framework. This will include descriptors for the material itself as well as for its interaction with the environment and elements to characterize intermediate effects in adverse outcome.
high-throughput screening (HTS) of bioactivity/toxicity is currently the only cost-efficient and rapid tool to screen the numerous nanomaterials (NMs) in use and under development. In the EPA ToxCast program, diverse classes of NMs and their micro-particle and ionic salt counterparts are tested in HTS assays to help prioritize NMs for further targeted testing. While the measured HTS endpoints are the same as those tested for traditional soluble chemicals, specific challenges arise for NMs for both experimental procedures and computational analysis. Challenges include no standard nomenclature, comparison of potency between different classes of NMs each tested at a different concentration range, linking NM physiochemical (pchem) characterization data into the analysis, and assay interference by NMs. With no standard nomenclature, we identified NMs by combining information on group (nano, micro, ion), chemical composition of the core and coating, primary particle size, and source. CAS numbers were purposely not used for NMs to avoid data being aggregated with bulk counterparts. We accounted NM exposure potential and tested NM at various concentrations, while all soluble chemicals were tested at the same concentrations in ToxCast and their sigmoidal AC50s were compared. NM bioactivity data is being analyzed using various dose metrics (e.g., either NM mass or surface area per medium volume or cell surface area) and potency estimates (LEC, sigmoidal AC50, etc.) for NM toxicity ranking. To link NM pchem properties into the bioactivity data, we choose to build a distinct database of NM pchem characterization results, instead of modifying the existing ToxCast database of bioactivity results. While HTS assays we used have successfully screened hundreds of soluble chemicals, inspecting the NM bioactivity results carefully resulted in discontinuing the use of one of the testing platforms because NMs interfere with the assay. Future research needs include developing computational models using NM pchem properties and/or in vitro data to predict in vivo effects.

**Computational Dosimetry for Nanomaterial Risk Assessment from Transcriptomic and HTP Data.**

L. G. Teeguarden, V. Mikhnev, J. G. Pounds, and B. D. Thrall. Systems Toxicology, Pacific Northwest National Lab, Richland, WA; Battelle Memorial Institute, Columbus, OH.

The convergence of evolutionary changes to the field of toxicology and the rapid emergence of an immense new class of toxicologically untested nanoscale materials has led to development of new genomic and high-throughput in vitro screening tools for toxicological assessment of nanomaterials. These advancements and refinements in the measurement and analysis of response have not however, been followed by equally important advancements nanoparticle kinetics and in the measurement of cellular or tissue exposure to test materials. The absence of adequate dosimetry data for nanomaterial toxicity studies is one of the largest sources of uncertainty in current testing paradigms. We present an integrated computational dosimetry framework for placing the results of in vitro genomics and high-throughput (HTP) toxicity data in the context of in vivo animal and human exposures and improving the basis for hazard rankings. The framework is applied in three case studies to demonstrate the impact of an evolution from “exposure” to target cell dose based nanotoxicity assessments: 1) the disruption of macrophage pathway clearance by macrophages in vitro; 2) in vitro cytotoxicity screening and hazard ranking of 25 metal oxides; and 3) in vitro screening and hazard ranking of nanomaterials from the U.S. EPA’s ToxCast program.

**Integrative Nanotoxicology: Linking Rapid Assays and Informatics to Predict Nanomaterial-Biological Interactions.**

S. Harper, Environmental and Molecular Toxicology, Oregon State University, Corvallis, OR; Chemical, Biological and Environmental Engineering, Oregon State University, Corvallis, OR.

While numerous nanotechnology and nanomaterial-based applications promise benefit to human health or the environment, the potential health and environmental risks associated with the unique properties of nanoscale materials are unknown and may lead to unintended health and safety consequences. The current gap in nanoparticle toxicological data dictates the need to develop rapid, relevant and efficient strategies to assess these risks, and may lead to unintended health and safety consequences. The current gap in nanomaterials and their behavior in the environment as well as their interactions with living systems. Data compiled using a dynamic whole animal (in vivo) assay to reveal whether a nanomaterial produces adverse responses has been made available through the Nanomaterial-Biological Interactions (NBI) knowledgebase (nbi.oregonstate.edu). Endpoints such as mortality, development, malformations and behavior were assessed in the embryonic zebrafish model. Data compression of the 25 individual endpoints provided the numerical representation of overall mortality and morbidity elicited by a particular nanomaterial at a given concentration. Computational analysis performed on 82 nanomaterial datasets has revealed the importance of surface chemistry as a driver for nanomaterial toxicity across material classes. Furthermore, data mining techniques to model the biological effects of nanomaterials were applied to this case study. Results illustrate that individual endpoints have different predictability given the same set of algorithms and cumulatively the combined values can provide significant insight into the potential toxicity associated with nanomaterials for use in hazard ranking.
(3) Outcomes on the JACVAM Comet validation and OECD guideline development, and (4) New approaches for genotoxicity assessments and guidance on dealing with positive results. In addition, this workshop will provide a forum to discuss the scientific advances that have led to the latest regulatory guideline changes, and their implementation.

W 793 US FDA Implementation of the ICH S2(R1) Guideline.
M. W. Powley, CDER, US FDA, Silver Spring, MD.

The genetic toxicology testing paradigm to support drug development was recently updated in ICH S2(R1) “Guidance on Genotoxicity Testing and Data Interpretation for Pharmaceuticals Intended for Human Use.” Although the guideline contained a variety of revisions, the most substantial impact was on the selection and conduct of genetic toxicity assays. Historically, in vitro and in vivo assays were performed to detect gene mutations as well as structural and numerical chromosome damage. While the updated ICH testing battery addresses these same endpoints in vitro, it also incorporates new in vivo endpoints. This update also included a variety of revisions, the most substantial impact was on the selection and conduct of genetic toxicity assays. Historically, in vitro and in vivo assays were performed to detect gene mutations as well as structural and numerical chromosome damage. While the revised ICH testing battery addresses these same endpoints in vitro, it also incorporates new in vivo endpoints. This update also included a variety of revisions, the most substantial impact was on the selection and conduct of genetic toxicity assays.

W 794 New Approaches in International Guidelines for Genetic Toxicology Assays: Latest Updates on OECD Guidelines.

In March 2010, the 22nd meeting of the Working Group of National Coordinators of the OECD Test Guidelines Programme (WNT) approved a project for updating the Test Guidelines on genotoxicity, with Canada, the Netherlands, France, and the USA identified as lead countries for this work. An Expert Working Group (EWG) comprised of international experts was established and has since met twice. The EWG has recommended the deletion of several Test Guidelines, including the as-conducted in yeast and Drosophila. The EWG is currently working on revisions to the in vitro Test Guidelines for the micronucleus test (TG477), chromosomal aberration test (TG473), and gene mutation test (TG474), as well as the in vivo assays for the mammalian erythrocyte micronucleus test (TG474), the mammalian bone marrow chromosomal aberration test (TG475), the dominant lethal test (TG478) and the spermatic gonosomal chromosomal aberration test (TG483). A new guideline dedicated to the thymidine kinase locus is also under development. In addition, an update has been initiated of the Introduction Document to the genetic toxicology test guidelines, which is intended to provide succinct and user-friendly guidance to guideline users. The purpose of the presentation at the SOT meeting will be to provide information on the status of the revision process and its main accomplishments to date. The opinions in this abstract are those of the authors and do not represent the policies of the U.S. EPA and FDA.

W 795 JACVAM Comet Validation and OECD Guideline.
M. Hayashi, Public Interest Incorporated Foundation, BioSafety Research Center, Shioh bridgen, Iwata, Japan.

In vivo rodent alkaline comet assay is used world-wide to detect genotoxicity of chemicals. The assay, however, has not been formally evaluated for its reliability and relevance. Thus, the Japanese Center for the Validation of Alternative Methods (JACVAM) has organized with ECVAM, ICGVAM, and NICEATM. This validation study has been supported by JEMS/SAMS and the consultation team including statisticians and specialists for the comet assay. We had the kick-off meeting for this trial in 2006. We have studied on the protocol optimization, intra- and inter-laboratory reproducibility before starting the predictive capability as the 4th phase. In the 4th phase-1st step validation study, the purpose was to examine the extent of reproducibility and variability of assay results among labs using cased test chemicals and the positive control EMS, when experiments were conducted in accordance with the Comet assay protocol-version 14. In review of the data, Validation Management Team (VMT) confirmed the reproducibility and variability of assay results among laboratories. Thus the VMT decided to move on to the 4th phase-2nd step validation study with an expanded set of test chemicals in accordance with the Comet assay protocol-version 14.2. The purpose of the 2nd step is to investigate the predictive capability of the assay against carcinogenicity of test chemicals. In the 2nd step, we selected 40 test chemicals, which include different characteristics in chemical classes including: Genotoxic carcinogen, Genotoxic non-carcinogen, Non-genotoxic carcinogen, and Non-carcinogen. Each test chemical was examined in one lab, because reproducibility was robustly confirmed in the 1st step validation study. We will disclose all data shortly but up to now VMT satisfied the outcomes. We are finalizing the validation report of our whole trial and we are also preparing the Comet Assay Atlas to guide how to select nuclei to be analyzed. Now, we are on the line of OECD TG through the peer review of our validation report and we have started to draft test guideline on in vivo comet assay.

W 796 New Approaches for Genotoxicity Assessment and Guidance on Dealing with Positive Results.
B. Gollapudi, V. Thibaud and J. Kim, ILSI-HESI IVGT Project Committee, Washington DC.

Exposure to mutagens can lead to adverse health consequences such as cancer and genetic diseases. A battery of short-term tests has been in use for decades to identify potential mutagenic agents. However, the relevance of findings from these assays in predicting human risk continues to be a subject of discussion, centering around three major topics on how to 1) improve the existing assays, 2) integrate emerging technologies, and 3) deal with positive findings. An international initiative consisting of experts from academia, industry and the government has been addressing these issues to enable more accurate assessment of human risk from exposure to mutagenic agents. In the area of improving the existing in vitro assays, the experts have been examining the choice of cell lines (e.g., human vs. rodents; p53 deficient vs. competent) and the metabolic activation systems (human vs. rodent) in order to make recommendations for future testing protocols. New technologies being considered as potential replacements and/or complements of the standard testing battery include the application of stem cells, 3-D skin/liver tissues, humanized animal models, high throughput assays (Tox21), incorporation of biomarkers of epigenetic changes with potential transgenerational significance, and the utilization of non-invasive imaging technologies to probe into time/age-related changes in the same animal with parallel anatomical and functional assessments. Results from the genotoxicity assays have traditionally been interpreted in a qualitative manner without analysis of the dose-response relationships. An expert group examined several quantitative approaches to the analysis of the dose-response data to identify a point of departure (POD) that could be combined with mode of action analysis to determine whether exposure(s) below a particular level (i.e., POD corrected by safety and uncertainty factors) constitute a significant risk for humans. These, along with other ongoing national and international initiatives are paving the way for a paradigm shift in the field of genetic toxicology.

PL 797 Early Markers for Screening Workers Exposed to Nephrotoxic Chemicals.
E. M. Metwally and A. ElSafy, Environmental & Occupational Medicine Department, National Research Center, Giza, Egypt.

Routine renal function tests are insensitive for detection of subclinical renal impairment. A marker of early renal affection is needed to be used for screening of workers at risk. In this study the activity of the urinary N-acetyl-B-D-glucosaminidase (NAG) was measured to detect early renal changes in workers exposed to nephrotoxic chemicals. The efficacy of NAG was also compared with that of urinary B2-microglobulin. The studied groups included 50 chemical laboratory workers exposed to aliphatic hydrocarbon solvents, 40 car painters exposed to aromatic hydrocarbon solvents, 36 plumbers exposed mostly to lead fumes and a control group of 36 clerks, matched the exposed groups in age, sex, smoking habits and socio-economic status.
Levels of urinary NAG differed significantly in all groups compared to controls. Unlike B2-microglobulin, NAG levels also expressed a strong and consistent correlation with cumulative exposure indices (HES,LES-Pb). Furthermore, an exposure-effect relationship existed between NAG and HES in solvent-exposed normotensive subjects. Values of NAG were higher in solvent-exposed hypertensives than normotensives. These results support the use of urinary NAG in the periodic screening of workers at risk, especially hypertensives and those with renal disease.

**Is Indoor Exposure to DEP a Health Risk?**

A. Gotti1, D.A. Sarigiannidou1,2 and S. P. Karakitsios1,3,4,5,6,7,8,9,10

The study describes a mechanistic approach for assessing the timecourse of the source-to-dose exposure to DEHP. The overall modelling framework was built on the assumption that DEHP is a common indoor air quality model, for estimating DEHP concentrations in the gas, particle, and dust phase starting from gaseous emissions of products containing DEHP.

- Multiphase indoor air quality model, for estimating DEHP concentrations in the gas, particle, and dust phase starting from gaseous emissions of products containing DEHP.
- Exposure assessment model that incorporates all possible exposure pathways and routes (inhalation of DEHP, skin exposure through dust rubbing off, non-dietary oral exposure through dust ingestion).
- Internal dose model, for the assessment of DEHP and its 3 major metabolites (MEHP, 5-0H MEHP and 5oxo-MEHP) in human tissues and urine through a multi-compartmental PBPK model.
- A variability analysis tool across all stages of the assessment.

Under a typical scenario in a common residential dwelling (surface area of 270 m2 and air exchange rate of 0.5 h-1) characterized by DEHP gaseous emissions of 200 μg/h (vinyl flooring and other plastic materials), the concentrations of DEHP in the gaseous, particle and dust phase are equal to 1.5 μg/m3, 21 μg/m3 and 4400 μg/kg settled dust. Overall daily intake varies between 0.2 to 10 μg/kg-bw, depending on the exposure scenarios considered. The latter are age-dependent: adults are exposed mostly through inhalation and infants through non-dietary ingestion. For a common repeated aggregate exposure scenario of 2 μg/kg-bw, the DEHP internal dose in venous blood and in adipose tissue (where bioaccumulation is clearly observed) reach a quasi-state equilibrium of 0.07 and 0.4 μg/gCr respectively. These findings are in good accordance to existing bio-monitoring data from NHANES. They are also one order of magnitude below the Biomonitoring Equivalent value of 660 μg/gCr for the sum of all DEHP metabolites measured in urine showing that DEHP concentrations in dwellings does not pose any significant risk.

**Identifying the Sources of Uncertainty in the Process of Reconstructing Exposures to Carbaryl Using Exposure-to-Dose Modeling.**

K. Holm1, R. McDougall2, M. Young3, B. Young1, H. J. Clewell4, R. Tornerovelez1, R. Goldsmith1, D. T. Chang1, C. M. Gruke1, M. B. Phillips1, C. C. Darby1 and C. Tan1

The research identified uncertainties in reconstructed carbaryl exposure estimates stemming from data gaps. Sources of uncertainty were identified by evaluating exposure reconstruction results in a simulated population for which we knew the exposure level. The “Cumulative and Aggregate Risk Evaluation System” (CARES) exposure model was used to generate time-concentration profiles for 500 virtual individuals exposed to carbaryl in food and drinking water for 365 days. Time-concentrations profiles were the inputs for a human physiologically based pharmacokinetic (PBPK) model for carbaryl, and biomarker (metabolite 1-naphthonol) concentrations in spot urine samples were simulated. Three mathematical techniques (namely, Exposure Conversion Factor, Discretized Bayesian, and Markov Chain Monte Carlo (MCMC)) were used to reconstruct mean daily carbaryl intake for the virtual population given the distribution of spot biomarker levels. We found that MCMC was the most precise method. Using MCMC for exposure reconstruction, we found that MCMC iterations did not converge to an exposure distribution if either the PBPK parameters or the urine flow rate were varied. Variation in the time between exposure events and spot sampling also contributed to the uncertainty in the reconstructed exposure. We recommend measuring urine flow rate, obtaining accurate estimates of PBPK model parameters, and monitoring time between exposure events and sample collection for improving reconstructed exposure estimates for rapidly metabolized compounds such as carbaryl.

**A Novel Design of Experiments Approach to Assess Biomarker Usage with Experiments Involving PBPK Models.**

J. Joy, M. A. Lyons, A. N. Mayeno, and B. Reifeld

In exposure assessment, samples from the environment and population are collected and analyzed to estimate the magnitude, frequency, and duration of exposure to an agent. In this and other applications, an appropriate design of experiments is critical to assure the optimal usage of resources and the efficient gathering of data necessary to answer important public health questions. To aid in such designs, physiologically based pharmacokinetic (PBPK) models are increasingly used to help elucidate chemical disposition, identify key biomarkers, and characterize the effects of uncertainty and variability. Although a number of approaches have been used to help optimize the design of experiments based on PBPK models, significant gaps remain in the efficiency and utility of these methods.

Here we describe a new approach in experimental design that can be used to identify quantities useful in exposure assessment experiments, such as maximally-informative biomarkers, types of biospecimen, and sampling times. This method involves the integration of an efficient sampling algorithm and metrics to assess information loss within a Bayesian inference framework containing a PBPK model. These features expand upon the capabilities of existing tools to include hierarchical statistical models and design comparison without random error.

To test the methodology, several models relevant to toxicology were evaluated, including those for styrene, dichloroethane, benzene, and chlorpyrifos. Results from these studies showed that the use of information loss as a design metric lead to precise and accurate reconstructions of simulated exposure data. These results also included a characterization and ranking of various biomarker- and exposure-related quantities in terms of their utility in conducting both forward and reverse dosimetry.

We expect that the proposed methodology will be useful in the design of exposure assessment experiments and related studies and will expand the field of Bayesian inference in environmental and human toxicology.

**Antioxidants and Oxidative Modification of Biomolecules: Can They Identify a Biomarker of Ozone Oxidative Stress?**

M. Kadiiska

The effect of ozone exposure on antioxidants and oxidation products of lipids, proteins and DNA in the plasma and urine of rats was studied by the international biomarker of oxidative stress study sponsored by NIEHS/NIH. The goal of this laboratory study was to identify a biomarker of ozone oxidative stress and to assess whether inconsistent results often found in the field of ozone oxidative stress might be due to a lack of comparability of the available methods where various oxidative products are measured as biomarkers. The time and dose-dependent effects of ozone exposure of rats on plasma and urine lipid hydroperoxides, TBA, malondialdehyde, isoprostanes, protein carbonyls, methionine sulfodioxiation, various tyrosine oxidation products, and DNA changes were investigated with different techniques. Effects of ozone on ascorbic acid, tocopherols, coenzyme Q, glutathione and DNA oxidation products, and DNA changes were investigated with different techniques. Ozone exposure did not cause statistically significant differences in plasma concentration products of lipid peroxidation or protein and DNA oxidation in a time- and dose-dependent pattern. However, urinary concentrations of isoprostanes measured with an immunoassay were increased by two different doses of ozone 8 h, 56 h and 70 h post-exposure. Since elevation of isoprostanes in urine was consistent at three time points studied, it is concluded that it fulfilled the oxidative stress biomarker criterion of significant effects measured in biological fluid and seen at both doses at more than one time point. Measurements of low molecular weight antioxidants in plasma are not sensitive biomarkers for oxidative damage induced by the ozone and may not be the tool of choice for the assessment of oxidative damage by ozone in vivo.
802 Enhancing PM Epidemiological Concentration-Response Functions by Incorporating Lung Deposition and Oxidative Potential.

S. P. Karakitsios1, D. A. Sarijanianni1, 2, V. Kalaitzis1 and M. Kermenidou1.
1Chemical Engineering, Aristotle University of Thessaloniki, Thessaloniki, Greece; 2Chemical Process and Energy Resources Institute, Centre for Research and Technology Hellas, Thermi, Greece.

Despite the fact that well documented associations have been established between urban air PM and mortality/morbidity, incorporating internal exposure and toxicity metrics would be expected to significantly improve risk assessment and management. Thus, existing concentration response functions were modified taking into account the fraction of particles deposited across the respiratory tract (using the MPPD lung deposition model) and the relevance to the corresponding health endpoint. The latter also depends on the actual size distribution for a given PM mass concentration, which is better described by the particle number count (PNC). Finally, the oxidative potential of particles of different size was also taken into account to derive revised exposure-response functions.

To investigate the feasibility of using this approach an extensive measurement campaign was carried out in a large Metropolitan area. PM size and number distributions were recorded in four sites. PM10, PM 2.5 and PM 1 samples were analyzed for oxidative potential by measuring Reactive Oxygen Species (ROS) using the DTT protocol. The results showed that the fraction of ultra-fine and fine particles is higher in the city center than in the suburbs. The same is true for oxidative potential especially for the smaller particles. Thus, the actual exposure for endpoints related to lower respiratory tract deposition and possibly translocation within the systemic circulation (e.g. cardiovascular disease, adverse pregnancy outcomes) might rise up to 4 times higher to the one estimated by the respective differences in mass concentration. Besides their implications for spatial epidemiology, our results show that future epidemiological studies would be greatly improved by incorporating evidence based on toxicity metrics, resulting in more robust associations between ambient air PM exposure and ill-health.


D. Close1, T. Xu2, J. Webb2, S. Ripp1 and G. Sayler1. 1490 BioTech Inc., Knoxville, TN; 2The University of Tennessee, Knoxville, TN.

Continuous toxicity monitoring using human cell-based bioreporters can report on the bioavailability and efficacy of toxic substances but is logistically challenging because of the cells’ relatively high autofluorescent background or the expense and/or sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similarly, sample destruction required to perform bioluminescent substrate (i.e., luciferin) addition. To facilitate this process, we have synthetically optimized the bacterial luminescence system (i.e., the pLUC100A plasmid) and the bioluminescent reporter cells were designed using a tetracycline activated promoter leading to a lower overall cost and increased data acquisition potential. Similar...
bile duct hyperplasia and hepatocellular hypertrophy in the PPARγ KO mice suggest a mechanism of hypertrophy independent of that previously reported for PPARγ-induced peroxisome proliferation in the current literature. Transmission electron microscopy (TEM) of selected liver sections from PND91 mice revealed PFOA-induced cellular damage and mitochondrial abnormalities with no evidence of peroxisome proliferation. These mitochondrial changes may represent PFOA-induced alterations in the regulation of mitochondrial permeability transition pores. This study was initiated by the NIEHS-funded BPA toxicity studies at the University of Michigan and other studies that have reported liver hypertrophy following perinatal exposure to BPA in rodents. These mitochondrial changes may represent PFOA-in-duced alterations in the regulation of mitochondrial permeability transition pores. This study was initiated by the NIEHS-funded BPA toxicity studies at the University of Michigan and other studies that have reported liver hypertrophy following perinatal exposure to BPA in rodents.

809 Fluopyram: Mechanistic Investigations to Elucidate the MoA for Liver Tumor Formation in the Rat.

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Fluopyram, a broad spectrum fungicide, caused hepatomegaly, liver hypertrophy and resulted in an increased incidence of hepatocellular carcinomas and adenomas in female Wistar rats following chronic exposure (24 months) at the highest dose evaluated (1500 ppm equating to 89 mg/kg/d) in the guideline carcinogenicity study. Mechanistic studies were subsequently conducted in the female rat to identify the initial key events responsible for the tumor formation and to establish thresholds for each of the early hepatic changes. Studies showed increased expression of constitutive androstane receptor (CAR) and pregnane X receptor (PXR) inducible genes from as early as 3 days of treatment. Further confirmation of CAR/PXR activation was provided by increased activity of specific Phase I enzymes (PROD and BROD respectively). Increased hepatocellular proliferation (measured by Ki67) was observed, particularly in the centrilobular region, starting from 3 days of treatment. Cell proliferation was also increased after 7 and 28 days of treatment but to a lesser extent than that following a 3 day exposure. In these studies, dose responses and clear thresholds were established for gene expression, enzyme activity and cell proliferation. Furthermore, these early hepatic changes were shown to be reversible following compound withdrawal. Other modes of action (MoA) for liver tumor formation such as genotoxicity or peroxisome proliferation were not observed. In conclusion, fluopyram is a threshold carcinogen and the resultant hepatocellular carcinomas in the female rat are due to hepatocellular proliferation mediated by CAR/PXR activation. It is unlikely that fluopyram would induce liver tumors in humans as the CAR/PXR induced hyperplastic response has been shown to be absent in human hepatocytes exposed to other known rodent hepatocarcinogens with the same MoA.

808 Liver Hypertrophy Is Not a Key Event in Constitutive Androstane Receptor (CAR)-Mediated Liver Tumor Development in Mice Treated with Triazoles.

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Liver hypertrophy is accepted to be a key event in CAR-mediated liver tumor development in rodents. Many triazoles (TRIs) have been reported to induce hypertrophy and tumors in the livers of rodents. In this study, we clarified the involvement of CAR in TRI-induced liver hypertrophy and tumor development using CAR knockout mice (CARKO). Seven-week-old male CARKO and wild-type mice (WT) were treated with cyproconazole (Cypro), tebuconazole (Teb), or fluconazole (Flu) for 2 weeks at 200, 150, or 200 ppm, respectively, in the diet. Tumors were initiated by a single i.p. injection of diethylnitrosamine (DEN) before the treatment. At interim kills at 4 (without DEN) and 13 weeks (with DEN), increases in liver weight and severe centrilobular/diffuse hepatocellular hypertrophy were detected in WT treated with TRIs. In CARKO, Teb induced severe liver hypertrophy, whereas Cypro and Flu induced very mild hypertrophy compared to that in corresponding WT. After 27 weeks of treatment, Cypro significantly increased eosinophilic altered foci/adenomas in WT. In CARKO, both types of lesions were clearly reduced compared to WT, but the multiplicity of foci was marginally higher than in the control. Teb increased eosinophilic foci in WT, but significantly decreased formation of these foci in CARKO. Flu and Teb had similar profiles in both genotypes. Basophilic foci/adenomas were also reduced in CARKO compared to WT for all TRI groups, but the effects were weak compared to changes in eosinophilic ones. Our present study indicated that CAR was the major mediator involved in liver hypertrophy for Cypro and Flu, but another pathway also existed. In particular, CAR was not involved in Teb-induced hypertrophy. Meanwhile, the involvement of CAR was crucial for TRI-induced tumors. These results suggested that liver hypertrophy was not a key event in CAR-mediated liver tumor development in mice.

810 Mechanistic Investigation of Technical Toxaphene (TT)-Induced Mouse Liver Tumors.

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Chronic exposure to technical toxaphene (TT) resulted in an increase in liver tumors in B6C3F1 mice, with male mice showing higher response. TT appears to act on the tumor promotion process. The current study was performed to further investigate the mode of action (MOA) of TT inducing mouse liver tumors. In a preliminary dose range-finding study, mice were given TT (0, 10, 40, 80, 160 and 320 ppm) in diet continuously for two weeks. A decrease in body weight gain compared to control was observed in mice treated with 320 ppm TT in diet (3.27 g vs. -1.22 g). Body weight gains were also significantly decreased in the 80 ppm and 160 ppm treatment groups. A significant dose dependent increase in relative liver weight was observed in mice treated with 0, 40, 80, 160 and 320 ppm TT diet (4.81, 5.32, 6.05, 6.55 and 7.43 g respectively). A similar trend was observed in the liver weight. A dose dependent increase in liver centrilobular hypertrophy was seen in mice treated with 80, 160 and 320 ppm TT; this was not observed at the 10 and 40 ppm dose groups. Similarly, a significant dose dependent increase in hepatocyte DNA synthesis (BrDU incorporation) was seen in mice treated with 40 ppm, 80 ppm, 160 ppm and 320 ppm, compared with the untreated controls with labeling indices of 3.28, 6.34, 11.10, and 15.61 respectively, compared with 1.06 in the control group). The increase in labeling index showed agreement with the absolute and relative liver weight results. Based on findings of the range-finding study, a second study using 0, 3, and 320 ppm TT diet was initiated to define the MOA for the TT mouse liver tumorigenic effects. Using phenobarbital as a positive control, the well-established hepatic tumor modes of action are being investigated including: cytoxicity; receptor activation (PPARγ, CAR, estrogen, and AhR) and oxidative stress in TT treated B6C3F1 mice.

811 In Vitro Cross-Species Comparative Analysis of Pharmacokinetic and Molecular Responses Mediated by a Labile AhR Activator.

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A novel herbicide in development, XDE-729 methyl, induces rodent liver effects through an aryl hydrocarbon receptor (AhR)-mediated mode-of-action (MoA). To further characterize species differences in these effects, mouse, rat, and human in vitro assays were used to elucidate differences in 1) AhR activation, 2) hydrolysis
rates to non-AhR activator metabolites, and 3) systemic exposure as predicted by a physiologically based pharmacokinetic (PBPK) model. AhR transactivation and binding assays indicated that XDE-729 methyl exhibits weak AhR agonism in mouse cells, no AhR agonism in human cells, and binds weakly to the rat AhR ligand binding domain. Fresh hepatocytes from CD-1 mouse, Sprague Dawley rat, and human (6 donor livers) were used to compare XDE-729 methyl mediated CYP1A1 induction, a sensitive biomarker for AhR activation. Gene expression analysis indicated that rats are the most sensitive species with regards to XDE-729 methyl mediated CYP1A1 induction with maximal induction at 100 μM of 147.7-fold in rat, versus 37.8-fold in mouse, and 2.9, 4.5, 6.2, 8.7, 10.3, and 34-fold in the six human livers. An in vitro study to determine hydrolisates rates of XDE-729 methyl to the major metabolites, XDE-729 acid, which does not activate AhR, indicated that: 1) the hydrolisate rate in human liver S9 is faster than rodents; 2) hydrolisate in human synthetic gastric fluid is faster than rodents; 3) hydrolisate of XDE-729 methyl is slowest in human whole blood versus rat or mouse. A PBPK model using the in vitro hydrolisate rates predicted similar blood and liver levels of XDE-729 methyl in rat and human after XDE-729 methyl dietary exposure. However, the in vivo mechanistic data indicate that humans are less sensitive than rats based on faster liver hydrolisate rates of XDE-729 methyl and limited AhR activation; therefore, a margin of exposure risk assessment using the in vivo rat toxicity studies is protective of human health.

812 Developmental Exposure to 2, 3, 7, 8 Tetrachlorodibenzo-p-dioxin (TCDD) Affects Leukemogenesis in Adult Tumor Prone Mice by Interacting with the Thymus Expressed Notch1.

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Reprogramming of progenitor cells during development can have profound impacts on later life disease susceptibility and is dependent on the interaction between genetic susceptibility in the embryo, the maternal environment during pregnancy, and the timing of exposure to potential insults. In particular, developmental exposure to the persistent contaminant TCDD, acting through the aryl hydrocarbon receptor (AhR), is known to cause immune suppression and is associated with hematological malignancies later in life. For example, at least 50% of all T-cell leukemias are associated with activating mutations in Notch1. Notch1 is a transmembrane receptor required for T-cell lineage commitment in the thymus. Moreover, preliminary data suggest a potential interaction between AhR and Notch1 signal transduction. To test for a possible gene-environment interaction between the AhR and Notch1, we hypothesized that leukemia prone Notch1 transgenic mice would have a more severe disease if exposed to TCDD during development. To test this hypothesis, timed pregnant dams were exposed to 0 to 3 μg/Kg TCDD throughout pregnancy. We found that while there was no difference in disease onset, severity or incidence, there was a significant difference in the cell-type of T-cell thymomas that developed in these mice and this difference was observed only in males. Specifically, whereas Notch-TG control mice developed CD4+ tumors, Notch-TG mice exposed to TCDD in utero developed CD8+ tumors. This lineage switch in disease phenotype suggests a reprogramming of a hematopoietic precursor or tumor stem cell during development leading to a reversal in the T-lineage choice in the adult life. These data have implications for disease susceptibility in vulnerable populations that may possess genetic instability in the Notch locus and/or have been exposed to AhR agonists development leading to a reversal in the T-lineage choice in the adult life. These data have implications for the Notch locus and/or have been exposed to AhR agonists development leading to a reversal in the T-lineage choice in the adult life.

813 Critical Evaluation of the Mode of Action of Carcinogenicity for Acrylonitrile.


Acrylonitrile (AN) is an aliphatic nitrile used as a reagent in industrial processes, and is polymerized into injection-molded plastics used to create pipes, automobile dashboards and children’s building blocks. Annually, over 1.5 million tons are produced in the U.S. alone, where AN is regulated by EPA as a hazardous air pollutant. Industrial cohort studies suggest a potential association between AN exposure and increased risk of cancer mortality and experimental evidence demonstrates that acrylonitrile induces tumor formation at multiple sites in rodent species following oral and inhalation exposure. A comprehensive cancer mode of action (MOA) analysis has been conducted and may inform the correspondence between the human and animal datasets. The AN literature was reviewed for evidence pertaining to genotoxicity and other potential MOAs; data were sorted by species, endpoint, and study design, and effects were critically evaluated for consistency, magnitude, specificity, plausibility and coherence. Early effects in relevant adverse outcome pathways were further assessed for causality, to determine if effects could indicate additional endpoints of AN carcinogenicity. Critical analysis of the AN data set supports potential contributions from multiple MOAs, including genotoxicity. Brain region-specific increases in lipid peroxidation and dose-dependent accumulation of 8-oxodG could support genotoxicity concurrent with oxidative stress in vivo, while covalent-binding to proteins and nucleic acids, liver DNA alklylation, leukocyte cell micronucleus formation and dose-dependent induction of forward mutations all support genotoxicity occurring prior to other cellular effects. While data-rich in some areas, other endpoints evaluated in the course of this analysis were data-poor. Some of these data gaps include a comprehensive evaluation of DNA repair and adducts, fixed mutations, and source of AN-induced oxidative species in rodent cancer target tissues. The views expressed are those of the authors and do not necessarily reflect the views or policies of the U.S. EPA.

814 Long-Term Inhalation Study with Nanomaterials: Pulmonary Effects of Nanoscale CeO2 and BaSO4 in a Rat 28-Day Range Finding Study.

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Inhalation exposure has been considered as the major route of concern for nanomaterials. Recent published literature reveals a distinct gap of long-term inhalation studies, especially on industrial relevant poorly soluble bio-persistent particles (PSP), including their carcinogenic potential. Nanoscale CeO2 and BaSO4 will be accurately examined in a combined chronic and carcinogenicity inhalation exposure study. Emphasis is placed on the relationship of inflammatory reactions, particle overload and lung tumor formation.

For this purpose, a 28 day range finding study according to the OECD Guideline 412 was performed. Groups of female Wistar rats (5 rats/group) were whole body exposed to 0.5, 5, 25 mg/m3 CeO2 and 50 mg/m3 BaSO4 for 28 days. A concurrent control group was exposed to clean air. The exposure concentrations for CeO2 were selected to achieve lung burden below and above particle overload. Biological effects were examined immediately after last exposure and in a post-exposure period. Examined endpoints included chronic and acute lung function, hematology and clinical chemistry, gross necropsy, histological examination of the respiratory tracts, and cell proliferation of the lung as well as systemic genotoxicity in peripheral blood cells. Increased lung weights in the high dose test group of CeO2 (25 mg/m3) and significant changes in lung lavage parameters (cell cytology, protein and enzyme levels) in the mid (5 mg/m3) and high dose test group of CeO2 are indicative for early treatment-related findings. Subsequent results of particle related biological effects associated with an appropriate particle lung burden of the range finding study will be presented here. The outcome of the 28 day study will serve as basis for concentration selection for the upcoming long-term inhalation study.

815 Pulmonary Responses in Rats after Inhalation Exposure to Cerium Oxide Nanoparticles Generated by the Harvard Versatile Engineered Nanomaterial Generating System (VENges).

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Current nanomaterials compounds have been used in a variety of consumer products including semiconductors, UV shields and diesel fuel additives to increase fuel combustion efficiency and decrease diesel soot emissions. Our previous studies have shown that exposure of rats to CeO2 by intratracheal instillation not only induces sustained pulmonary inflammation, but also lung fibrosis. In the present study, the aerosols of CeO2 or CeO2 coated with a nanothin layer of amorphous SiO2 (aSiO2-CeO2) were generated by the Harvard VENGES. The aerosols were diluted with air and delivered to a whole body exposure chamber. Male Sprague Dawley rats were exposed to CeO2 or aSiO2-CeO2 at 2.7 mg/m3, 2h/day for 4 days along with air controls. Animals were sacrificed at 1or 84 days post exposure. Morphometric analysis of the CeO2 and aSiO2-CeO2 particle cores showed diameters of 12.8 and 2.4 nm, respectively. Morphology diameter modes of 82 and 96 nm were measured for the CeO2 and aSiO2-CeO2, aerosols aggregates in the breathing zone of the animals. Alveolar macrophages (AM) were obtained by bronchoalveolar lavage (BAL), and acellular BAL fluids (BALF) were saved for further analysis. At 1 day after CeO2 exposure, but not aSiO2-CeO2 significantly induced PMN infiltration and lactate dehydrogenase activity in the BALF. CeO2 significantly increased collagen
degradation enzymes, matrix metalloproteases (MMPs)-2 and tissue inhibitor of MMP-1 in the BALF, which may be involved in the modification of the extracellular matrix. At 84 days post exposure, none of the particle treatment groups induced lung inflammation, cellular injury or alteration of hydroxyproline content in lung tissues. These results demonstrated that a thin coating of aSiO2 on CeO2 protected lungs from CeO2-induced acute lung toxicity, suggesting that a thin coating of aSiO2 may potentially be used to modify other nanoparticle-induced lung toxicity.

**Gene Expression Profiling of Human Lung Epithelial Cell Lines Exposed to Manufactured CoO and CeO2 Nanoparticles.**

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Exposure to manufactured nanoparticles (NPs) via inhalation can cause adverse human health effects. A transcriptomics study was performed to identify molecules and cellular pathways that are specifically triggered by in vitro exposure of the human bronchial BEAS-2B and alveolar A549 epithelial cell lines to 7-nm CoO and 4-nm CeO2 NPs. We aimed to investigate whether 1) the same lung cell type responds similarly to the NPs; 2) alveolar vs. bronchial epithelial cells respond differently to the same NP; and 3) immunological processes are influenced. Non-cytotoxic exposure concentrations of monodispersed NPs were used. Statistically significant changes in gene expression as compared to solvent-treated cells (median fold-change>1.5, p<0.05) were evaluated after 3, 6, 10, and 24 hours. The kinetics of the cell responses induced by the 2 NPs were similar within, but different between the 2 cell models. BEAS-2B cells were found to be more sensitive for NP toxicity, as they showed a higher total number of differentially expressed transcripts (DET) at a 10-fold lower NP-concentration than A549 cells. Hierarchical clustering of all DET indicated that the transcriptional responses were quite heterogeneous among the 2 cell types and 2 NPs. Between 1% and 14% DET encoding markers involved in immune system processes were observed in the BEAS-2B and A549 cell lines, resp., with the highest fractions observed in BEAS-2B cells. Most of these genes, i.e., ITGB2, TLR6, PAG1, HLA-DRB3, TIRAP, and HLA-A, are involved in immune signalling or yet unassigned pathways. Nanoparticle exposure mainly induced suppression of immune gene transcription, rather than immune stimulation. The AKT1 gene was identified as a possible generic marker of lung epithelial cell-NP interaction. Our data suggest that CoO- and CeO2-NP give rise to a distinct immunological response in bronchial and alveolar epithelial cells.

**Mechanistic Insights into the Toxicity of Multivalved Carbon Nanotubes and Cerium Dioxide Nanoparticles in Primary Human Bronchial Epithelial Cells.**

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Cerium dioxide nanoparticles (CeO2 NPs) and multi-walled carbon nanotubes (MWCNT) are priority materials for urban risk assessment due to wide spread industrial, consumer product and environmental utilizations. We aimed at deciphering the impact of CeO2 NPs and MWCNTs on primary human bronchial epithelial cells (BEC) following an ex-vivo exposure. CeO2 NPs and MWCNT suspensions were thoroughly characterized, including using transmission electron microscopy (TEM), dynamic light scattering (DLS), and zeta potential analysis. Cells were then exposed to nanomaterials for 18-24 hours and mechanisms of cell injury were studied. TEM revealed that both CeO2 NPs and MWCNTs are internalized by bronchial epithelial cells and are found either in vesicles or free in the cytoplasm. CeO2 NPs fail to elicit a toxic response in BECs at environmentally relevant doses. However, diesel exhaust particles and CeO2 NPs co-exposure leads to significant increase in cytotoxicity. MWCNT exposure in bronchial epithelial cells leads to a time-dependent decrease in viability, increased reactive oxygen species production, NF-kB (p65)/Rel A phosphorylation and nuclear translocation. Moreover, we observed caspase-1 activation and increased numbers of autophagic vesicles in MWCNT-treated cells as compared to control cells. An increase in p62 levels indicated a block in autophagic turnover rather than autophagy induction in these cells which was associated with cytoskeletal alterations induced by MWCNT. In conclusion we demonstrate that nanomaterials exposure lead to toxic events in primary human bronchial cells. Moreover, we showed that doses of CeO2 NPs and diesel exhaust particles that are innocuous in themselves can result in toxicity when given as a co-exposure.

**Inflammatory and Free Radical Generation Characteristics of Nano-Cerium Dioxide.**

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Nano-cerium dioxide (CeO2) possesses the potential for use in human health by protecting against the deleterious effects of ischemia and radiation. However, the literature is polarized about the effects this compound has in vivo. Our laboratory has shown that pulmonary nano-CeO2 impairs arteriolar reactivity 24 hrs post-exposure. The mechanisms of this impairment are currently unknown but may be linked to the free radical scavenging or inflammatory properties of this nanoparticle. The aim of this study was to: 1) thoroughly assess the physical and chemical characteristics of the nano-CeO2; 2) examine the antioxidant potential of nano-CeO2 via electron spin resonance (ESR); and 3) assess the pulmonary inflammation in Sprague-Dawley rats 24 hrs post-intratracheal nano-CeO2 instillation. The primary particle size of the nano-CeO2 was calculated to ~3 nm (via transmission electron microscopy and surface area measurements). Dynamic light scattering determined the agglomerate size (~80 nm) and x-ray photoelectron spectroscopy determined the valence state of the nano-CeO2. The ESR measurements indicated that nano-CeO2 alone did not generate free radicals and in the presence of cells (Raw264.7), nano-CeO2 quenched the free radicals generated by these cells. Finally, bronchial alveolar lavage from rats instilled with 0, 10, 100 or 400 µg of nano-CeO2 revealed an increase in polymorphonuclear leukocytes (0.6±1.2, 0.8±0.3, 7.1±0.9, and 10.3±0.9 per 106 cells), and lactate dehydrogenase (90±12, 100±9, 453±53, and 60±2±0.2 units/L) but there was no change in albumin. These findings provide evidence that pulmonary inflammation is present after exposure but does not damage to the epithelial/endothelial cellular barrier. Additionally, these nanoparticles are capable of quenching free radicals there by exerting a systemic effect.

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**Nanoceria Distribution and Biopersistence in Rats Is Not Consistently Affected by Particle Size, Shape, Dose, or Dosing Schedule.**

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Background: Nanoceria is a diesel fuel additive, an abrasive in integrated circuit fabrication, and is being developed as an antioxidant therapeutic. Objectives: Determine the influence of nanoceria size, shape, dose, and dosing schedule on its distribution and biopersistence. Methods: Aqueous dispersions of citrate-stabilized cubic or polyhedral ceria and a ceria nanorod (10 x 40 to 600 nm), synthesized and characterized in-house, were iv infused into rats (single infusion of 5 nm @ 11, 56 or 85 mg/kg, 15 nm @ 70 mg/kg, 30 nm @ 6 or 85 mg/kg, 55 nm @ 50 mg/kg, nanorod @ 20 or 50 mg/kg; and 5 nm @ 11 mg/kg for 30 consecutive days). They were terminated 1 h to 90 days later. Controls received vehicle. Multiple organs were weighed and samples collected from multiple sites and blood for cerium determination. Results: The greatest % of the dose was in the liver, spleen and bone marrow; these levels decreased over time only in liver for the 30 nm ceria @ 6 mg/kg, and increased in spleen and bone marrow over time in several cases. There were no consistently significant differences in the % of the dose in the liver or spleen for the different sizes, shapes, doses, or dosing schedules other than tendencies for more nanorod accumulation in the spleen than the 5 nm polyhedral ceria and more nanorod accumulation in the bone marrow than the 5 or 30 nm ceria. Brain nanoceria was low; little to none was in brain parenchyma. Conclusions: Nanoceria, an insoluble metal oxide, was cleared into mononuclear phagocyte system organs in which it persisted for 90 days. Size, shape, dose, and dosing schedule had little effect on its distribution or persistence, suggesting repeated exposure will likely produce accumulation, perhaps reaching a level shown to be toxic after single high-dose administration. Support: US EPA STAR Grant RD-833772.
Metal Oxide Nanoparticles After Spontaneous Activity and Pharmacological Responses in Neuronal Networks Grown on Microelectrode Arrays.

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The widespread use of engineered nanoparticles (NP) has increased their exposure potential and made it necessary to assess potential impacts on human health. Previous studies indicate that NPs can enter the brain via the olfactory nerve or by crossing the blood-brain barrier; thus, it is essential that their effects on neuronal function be examined. In the present study, 5 CeO2 (7 to 1388 nm), and 4 TiO2 (6 to 200 nm) NPs were examined to determine their ability to alter network function in primary cultures of cortical neurons grown on 12 well microelectrode array (MEAs) plates. NPs were dispersed in Neurobasal A medium containing 20% FBS (dispersant). Between days 14 to 21 in vitro, 1 hr of baseline activity was recorded prior to exposure to NPs. Changes in spontaneous mean firing rate (MFR) relative to the dispersant control were assessed 1, 24, and 48 hrs after exposure to NPs (3–50 μg/ml). Following the 48 hr recording, the response to a pharmacological challenge with the GABAβ antagonist bicuculline (BIC; 25 μM) was assessed. In all, 3 of 5 CeO2 and 2 of 4 TiO2 NPs decreased MFR below threshold following 1 hr exposure. All 4 TiO2 NPs altered MFR beyond threshold at 24 hrs. At 48 hr, the MFR for all 9 NPs deviated from minimum threshold. BIC increased MFR in dispersant treated networks; of the 9 NPs tested, 3 TiO2 particles and 1288 nm CeO2 increased the BIC response in MFR, while <7 nm CeO2 suppressed the BIC-induced change in MFR relative to control. Most notably, the results show that changes in network function occur in the absence of cytotoxicity and were observed at all time points regardless of particle type. The results indicate that metal oxide NPs can disrupt both spontaneous and GABAβ receptor-mediated neuronal activity in vitro. Additional studies are necessary to investigate the mechanisms underlying these observations and understand the implications of NP exposure to neuronal function in vivo. (This abstract does not reflect Agency Policy.)

Network2Canvas: Network Visualization on a Canvas with Enrichment Analysis.

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Networks are vital to computational systems biology research, but visualizing them can be a challenge. For networks larger than ~100 nodes and ~200 links, ball-and-stick diagrams fail to convey much information. To address this, we developed Network2Canvas (N2C), a web application that provides an alternative way to view networks. N2C visualizes network nodes by placing them on a square toroidal grid. These nodes are then clustered together on the grid using simulated annealing to maximize local connections. For visualizing the canvas, a node’s brightness is made proportional to its local fitness; the brighter a node is, the stronger its connections to its neighbors. The grid is interactive, implemented in HTML5 and the JavaScript library D3. We applied N2C to create canvases for 25 gene-gene functional association networks connecting human and mouse genes, and six drug-drug network connections FDA approved drugs based on their shared properties. N2C also has functions to perform enrichment analysis. Given lists of genes or drugs, N2C highlights enriched terms on the grid as well as computes the degree of clustering for these enriched nodes. We applied N2C to analyze nine cancer cell lines by comparing their enrichment signatures to enrichment signatures of matched normal tissues. Such analysis provides a global visualization of critical differences between normal tissues and cancer cell lines. In particular, we observed a common pattern of up regulation of the polyclonal group and enrichment for the histone mark H3K27me3 in many of cancer cell lines. In summary, N2C provides a new flexible method to visualize large networks and to perform and visualize enrichment analysis on functional drug and gene networks. N2C is freely available at http://www.mayanlab.net/N2C

Computational Approaches to Predicting Adverse Drug Reactions and Mitigating Off-Target Liabilities in Early-Stage Drug Discovery.

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Identifying unintended off-targets associated with adverse drug reactions (ADRs) is daunting by empirical methods alone. Many of these are unrelated to the therapeutic target and it would need hundreds of assays to reveal their involvement in clinical side effects. However, latest development in in vitro assay technologies, tools for pharmacovigilance, and establishment of clinical data warehouses laid the foundation for the development of computational strategies to predict side-effect targets and even their associations with pathways and complex biological systems. However, many of the early methods focused on chemoinformatics aspects of lead optimization with little attention for translational value, not to mention the safety aspects of the chemical structures and individual molecules. Introduction of several theoretical methods to explore drug-target-adverse reactions changed the landscape of safety assessment and the translational value of these new methods has been increasing significantly. The methods could be characterized as statistical models, expert or rule-based systems, quantum mechanics calculations, and structure based approaches. Data used by these models are largely generated in in vitro assays and obtained from clinical observations. Novartis, in collaboration with UCSF and SeaChange has conducted a large-scale experimental test of the SEA method to predict drug activities on targets linked to ADRs, and a guilt-by-association method to associate that activity with side-effects and provided evidence that a drug-target-ADR network approach could have widespread application to de-risking toxicological liabilities in drug discovery. In this presentation, we will highlight the various methods and applications developed for early risk assessment, and provide insights of their practical applications.
Because of their complexity, cutting-edge machine-learning methods will be critical for building systems models of cell and tissue behavior and for future drug development. Such models require accurate information about the subcellular distributions of proteins, RNAs and other macromolecules in order to be able to capture and simulate their spatiotemporal dynamics. Unfortunately, information with sufficient resolution and for different cell and tissue types is currently very limited. Microscope images provide the best source of this information, and new tools are being developed to build models of cell organization directly from such images. The number of possible experiments needed is prohibitive, but active machine learning methods can be used to help choose which experiments are needed in order to construct sufficient models of how cell organization changes with cell type and with disease. In order to use these models for drug development, information on how thousands of cell components respond to millions of potential therapeutics is also needed in order to minimize toxicity. Active-learning methods can also guide experimentation to overcome the dimensionality of this problem.

The adverse effects of air pollution on cardiovascular health have been established in a series of major observational studies. Even brief exposures to air pollution have been associated with marked increases in cardiovascular morbidity and deaths from myocardial ischemia, arrhythmia, and heart failure. The breadth, strength, and consistency of the evidence provide a compelling argument that air pollution, especially traffic-derived pollution, causes cardiovascular disease. However, these observational data are limited by imprecision in the measurement of pollution exposure, and the potential for environmental and social factors to confound these apparent associations. For a causal association to have scientific credence, a clear mechanism must be defined. What are the potential pathways through which air pollution mediates these adverse cardiovascular effects and diseases? And are the effects caused by the nano-sized particles? This session will focus on the underlying biological mechanisms of complex particle mixtures.

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Fine particulate matter (PM2.5), gaseous co-pollutant, and noise exposure were assessed in a panel of workers who spent most of their professional activities on or near roads. To quantify workers’ exposure and related short term health effects, up until now, we observed 13 road maintenance workers, each over 5 non-consecutive work days. We used a methodology based on personal and work site measurements to assess the workers’ exposure to particles, noise and co-pollutants. For examination of personal exposures during and after work, the workers were equipped with a personal particulate monitor (PM2.5) and a noise dosimeter. Additional exposure parameters measured at the work site provided a detailed evaluation of exposure during work. These included ultrafine particle counts, measurement of carbon monoxide, nitrogen dioxide and ozone as well as PM sampling for compositional analysis. Cardiovascular health endpoints were assessed before, during and 15 hours post work shift. These included a continuous ECG from before until 15 hours post work, measurement of blood pressure and lung function before and 15 hours post work as well as measurement of exhaled nitric oxide and blood markers 15 hours post work shift. Noise levels are generally high with extreme levels during certain activities. Mean work shift PM2.5 Mass concentrations ranged from 17.0 μg/m3 to 421.0 μg/m3 (mean 56.8±17 μg/m3) and UFP counts were between 15.338 particles/cm3 and 408,518 particles/cm3 (70,721 particles/cm3). Results suggest that PM2.5 induces cardiovascular effects and inflammation. PM2.5 and noise were only weakly correlated, allowing further assessment of these associations.

Traffic-generated pollutant-exposures appear to have a strong correlation to these adverse vascular outcomes, as shown through roadway proximity studies. Controlled toxicological studies highlight potential interactions between vehicle-source emissions and upregulation of signaling molecules associated with progression of atherosclerosis. Mechanistically, a role for both innate and adaptive immune responses is emerging, with important recent findings demonstrating that receptors such as the lectin-like oxidized LDL receptor (LOX)-1 may play a role in communicating airway exposures to cardiovascular outcomes. Using a mixed vehicle emission (MVE; gasoline and diesel engine, 100 PM μg/m3) model we show that atherogenic Apolipoprotein E null (Apoe-/-) exhibit increased oxidized LDL, associated with proinflammatory responses and lipid accumulation, as well increased vascular LOX-1 expression, when exposed by inhalation 6 h/d for 7 days, compared to filtered air-exposed controls. Furthermore, we observe a significant upregulation of vascular factors downstream of the LOX-1 receptor that are associated with atherosclerotic plaque growth and rupture, including reactive oxygen species, endothelin (ET)-1 and matrix metalloproteinase (MMP)-9 in MVE-exposed Apo E-/- mice, expression of which are attenuated with anti-LOX-1 antibody treatment. These data indicate that vascular effects of inhalation exposure to traffic-generated pollutants, resulting in progression of atherosclerosis and onset of clinical cardiovascular events, may be mediated through scavenger receptor ligand binding, internalization, and downstream signaling pathways.

Explication of a mechanism lends substantial scientific credence to epidemiologic associations. For example, there are strong epidemiologic associations between increases in air pollution, specifically traffic-related air pollution, and adverse cardiopulmonary outcomes, but our understanding of explanatory mechanistic pathways is incomplete. What are the potential pathways through which air pollution mediates these adverse cardiopulmonary effects? This presentation will focus on some of the underlying biological mechanisms. One specifically hypothesized mechanism is acutely increased oxidative stress/inflammation in the airways, as reflected in exhaled breath condensate (EBC) nitrite and nitrate in human subjects. Nitric oxide (NO) is formed in the lung epithelium from activities of all three NO synthases, and may then diffuse into the airway lining fluid where it is subject to oxidation to nitrite and nitrate by multiple pathways. Data showing an increase in exhaled breath condensate nitrite and nitrate following either controlled diesel exhaust inhalation or acute highway traffic exposure will be reviewed. Multiple oxidative pathways for the acute increases in EBC nitrite will be discussed, along with implications for generation of cardiopulmonary effects.
metric for oxidative stress in vivo could be the oxidative potential of particles indicating the possibility of PM to produce oxygen radicals and cause damage in vivo. The focus of this presentation will be on the oxidative potential of PM from different sources as a measure to predict source specific cardiovascular toxicity in vivo. Systemic effects in rats exposed for 4-weeks to diesel engine exhaust occurred including decreased numbers of white blood cells and reduced von Willebrand factor in the circulation. In addition, lung tissue factor activity is reduced in conjunction with reduced lung tissue thrombin generation. We have also shown in studies in which volunteers were exposed to air pollution at various locations dominated by different sources of emission that organic carbon, nitrate and sulfate to be most consistently linked with different biomarkers (e.g. high-sensitivity C-reactive protein, fibrinogen, von Willebrand factor and plasminogen activator inhibitor-1) of acute cardiovascular risk. Associations for PM mass concentrations and OP were less consistent, while other measured components of the air pollution


The detrimental effects of air pollution on the cardiovascular system are now well-established, however, the mechanisms underlying these effects remain to be determined. Using diesel exhaust as an example of a common air pollutant, rich in combustion-derived nanoparticles, this presentation will give an overview of the multiple ways urban air pollution can alter cardiovascular function. Studies in both animals and man have demonstrated that this pollutant impairs vascular function, promotes thrombosis, exacerbates cardiac ischemia and accelerates the development of atherosclerosis. The presentation will focus on preclinical experiments with diesel exhaust particulate (DEP) that provide insight into potential mechanisms for these effects, including oxidative stress, endothelial dysfunction, inflammation, particle translocation, as well as considering with constituents of DEP may be responsible for these harmful effects.

832 The Dynamics of Neuroinflammation and Inflammatory Cell Responses in Neurologic Disease.
G. J. Harry1 and C. P. Curran.1 NTP Laboratory, NIEHS, Research Triangle Park, NC; 2Northern Kentucky University, Highland Heights, KY.

An increasing body of evidence indicates that neuroinflammation and activation of immune cells within the nervous system are associated with neurodegenerative disease, neurodevelopmental disorders, and potentially in reaction to environmental exposures. However, it is also increasingly obvious that these responses may be beneficial or detrimental and discriminating between these is now only recently been addressed. Understanding the process by which these responses are triggered and the spatiotemporal dynamics of the response is critical to developing a strategy for translating neuroinflammation and immune response to effective prevention or treatment of neurologic disease/injury. Three well-known neurotoxicants: trimethyltin, manganese, and mercury will be used as chemical probes to explore differences in the timing of inflammatory effects and consequences in the brain. We will conclude by discussing approaches to manipulate the lesion microenvironment and/or brain macrophage such that inflammation favors tissue repair in the spinal cord.

833 Microglia Heterogeneity in Neuroinflammation and Neurotoxicity.
G. J. Harry. NTP Laboratory, NIEHS, Research Triangle Park, NC. Sponsor: C. Curran.

Microglia cells are the resident immune cells of the brain; however, they are also critical neural specific cells with multiple roles. The response of microglia has been associated with environmental PM exposures and is present in diagnosed neurodegenerative disease. In many cases, what is considered a microglia response is often a brain macrophage response that can be derived from both resident microglia and infiltrating blood-borne monocytes. In order to better understand the resident microglia response, data will be presented from a delayed neuronal death model following trimethyltin intoxication. This compound induces death of the dentate granule neurons across multiple species and is accompanied by a robust microglia response and elevation in pro-inflammatory cytokines. The process is tightly regulated and with the phagocytic clearance of the neuronal debris, the system actively undertakes repair. Using this model of resident microglia activation, the dynamic sequence of events will be characterized and the role of the microglia response within areas of neuronal death, synaptic loss, and neuronal activity. Using this model we are able to identify the various functions of microglia and potentially identify a profile of molecular and morphological responses that will identify the different functions of the microglia. This will serve to identify those processes of microglia that need to be fostered and those that need to be mitigated by therapeutic intervention.

834 Neuroinflammation and Developmental Vulnerability to Manganese.
R. B. Talbun1, 2, 1Center for Environmental Medicine, Colorado State University, Fort Collins, CO; 2Toxicology Section, Department of Environmental and Radiological Health Sciences, Colorado State University, Fort Collins, CO.

There is increasing evidence that activation of microglia and astrocytes during development can influence susceptibility to neurodegeneration later in life. Environmental insults such as infectious agents, pesticides, and heavy metals have all been implicated in neuronal injury leading to increased activation of glial cells and development of chronic neuroinflammation. Exposure to elevated levels of the essential element manganese (Mn) causes a spectrum of neurochemical and neuropathologic changes that can culminate in irreversible neuronal injury in subcortical and cortical structures. Children are more vulnerable to Mn than adults and recent epidemiological evidence links high Mn in drinking water to cognitive and behavioral impairment. Mn neurotoxicity is associated with astrogliosis in the basal ganglia and studies conducted in our laboratory and others suggests that glial-derived inflammatory cytokines and nitric oxide (NO) influence the progression of neuronal injury. Increased expression of iNOS/NO2 by activated glial cells in response to Mn results in nitrosative stress throughout the basal ganglia and enhanced apoptosis within selected populations of neurons in the globus pallidus and striatum. NO2 is exclusively expressed in glia and produces high levels of NO, which forms highly reactive peroxynitrite anion (ONOO-) upon combining with superoxide, resulting in electrophoric nitration of cellular proteins that damages neurons. Peroxynitrite-mediated nitrosative stress is also implicated in a number of neurological disorders, including Alzheimer’s and Parkinson’s diseases. Inflammatory changes in glial cells may therefore be an important link between neurotoxic injury and persistent neurological dysfunction during aging.

835 Spontaneous and Mercury-Induced Antibodies to Brain Antigens Affect Fetal Brain Development.
D. A. Lawrence1, Y. Zhang1, Y. Hsu2, and D. Gao.1 Immunology, Wadsworth Center, Albany, NY; 2Occupational Health, Catholic University of Daegu, Daegu, Republic of Korea.

BTBR mice spontaneously develop a high level of serum IgG of which a proportion are antibodies (Abs) to brain antigens. These Abs enter the brain and are associated with expression of inflammatory cytokines, which are suggested to be related to presence of activated microglia and mast cells. The neurodevelopmental effects of BTBR mice are posited to be autoimmune due to elevated presence of activated CD4+ T cells driving Abs production, and the offspring have behaviors that resemble autism. IgG from BTBR mice intravenously injected into B6 pregnant dams caused offspring to have lowered social interactions; likewise, B6 embryos that are born from BTBR dams have lowered sociability. Changes in mitochondrial functions can alter these behavioral outcomes of BTBR mice, but the mechanisms remain unknown. Abs causing behavioral deficits is not a new finding, in that Abs induce neuropsychiatric syndromes in lupus-prone mice. Additionally, certain strains such as A.SW, which are sensitive to Hg-induced autoimmunne responses, generate autoantibodies to brain antigens upon developmental exposure to HgCl2 and have behavioral deficits as adults, and inflammatory cytokines in multiple brain regions are associated with the behavior changes; however, the HgCl2-induced effects are dependent on the genetics of the strain and sex of the developmentally exposed mice.

836 Manipulating Microglia and Macrophages to Promote Repair of Injured Spinal Cord.
P. G. Popovich. Department of Neuroscience, The Ohio State University, Columbus, OH. Sponsor: C. Curran.

Following traumatic or ischemic/reperfusion injury to spinal cord or brain, macrophages derived from resident microglia and infiltrating blood monocytes accumulate in the affected area. Collectively, these cells exert diverse and conflicting effects on neurons and glia. Indeed, microglia/macrophages can cause neuronal cell death, axonal injury and demyelination but can also promote neuron survival, axon regeneration, remyelination and revascularization. Over the past few years, we have
used a number of strategies to manipulate resident and recruited macrophages in an effort to understand their seemingly paradoxical functions, specifically with consideration of their role in tissue repair vs. cell killing. This study demonstrates that resident and recruited macrophages exhibit distinct pro-inflammatory and anti-inflammatory phenotypes, and that their differentiation can be manipulated with specific cytokines and chemokines.

### 837 Advances in Carcinogenic Risk Assessment of Low-Level Genotoxic Impurities in Pharmaceuticals.


Pharmaceutical syntheses involve the use of reactive starting materials, intermediates, and reagents, some of which are known or potential genotoxicants and carcinogens. Therefore, genotoxic and potentially carcinogenic impurities may appear in the final drug product. Risk assessment approaches have focused on defining impurity limits which pose acceptable risk over a patient’s duration of drug treatment. Over the past decade, regulatory guidances (EMA, US FDA draft) have been introduced. Drug-associated genotoxic impurities became an ICH guideline topic (M7) in 2009, and an Expert Working Group is currently developing a harmonized guideline. The ICH M7 effort presents an opportunity to review existing guidelines, evaluate new information and experiences since their introduction, and improve the integration of safety and quality aspects for detection, risk management and control. This workshop will provide a historical overview and introduce newer concepts to SOT members for discussion on several approaches being considered in M7. The workshop will cover the following topics along with case studies: (1) A brief overview of regulatory risk assessment approaches developed over the past decade and introduce current M7 concepts being considered; (2) Review current in silico Q(SAR) and genotoxicity testing approaches to predict or identify hazards and quality impurity risk; (3) Introduce advances in the framework and rationale for applying next generation acceptable risk limits during clinical development and marketing; (4) Highlight differences in chemical space between pharmaceutical synthetic intermediates or impurities and that used to derive the original lifetime threshold of toxicological concern (TTC) limit, and its potential implications for risk characterization; and (5) Review approaches for addressing risk of multiple genotoxic impurities in a drug product.

### 838 Genotoxic Impurities—Regulatory Advances in Risk Assessment Approaches.

P. Kasper. Federal Institute for Drugs and Medical Devices, Bonn, Germany. Sponsor: W. Ku.

The control of impurities in drug substances/products is regulated by internationally harmonized guidelines (ICH Q3A/B). However, these documents lack any specific instructions on how to treat impurities with a genotoxic potential. This has led to considerable differences between regulatory authorities on what levels of daily intake of genotoxic impurities are acceptable and prompted the European Medicines Agency (EMA) to release a “Guideline on the Limits of Genotoxic Impurities” in 2007 followed by publication of a draft guidance by CDEFA in 2009. Several basic principles are common to both documents and are currently under discussion for further development in the ICH process.

Genotoxic impurities are defined as compounds that are DNA-reactive and have the potential to directly cause DNA damage when present at low levels leading to mutations and therefore, potentially causing cancer. Consequently, in silico prediction of DNA reactivity/mutagenicity and Ames mutagenicity testing are basically the recommended approaches for hazard identification. Since risk assessment based on such limited set of data is not feasible a generic “Threshold of Toxicological Concern” (TTC)-value of 1.5 μg/day corresponding to a 10-5 lifetime risk of cancer is used as a pragmatic approach for control of impurities that are Ames-positive. Derived from linear extrapolation of rodent potency data from over 700 carcinogens and based on an accumulation of worst-case assumptions the TTC value is a very conservative and therefore sufficiently safe limit. Possible deviations from the default TTC approach are currently under discussion and include cases when more extensive and appropriate data for a proper assessment of potential risks are available, e.g., data from carcinogenicity studies with the impurity or mechanistic data providing evidence for a threshold. Also specific clinical aspects may justify an adjustment of the default data level of genotoxic impurities, such as for instance treatment of life-threatening condition, short life expectancy of patients, or indications with less-than-lifetime exposure.

### 839 Advances in Mutagenic Impurity Hazard Assessment: Best Practices and Current Developments.


For many years, pharmaceutical companies have been using in silico methods as a primary tool for the identification of mutagenic impurities or degradants in drug substances or products as recommended in the European Medicines Agency (EMA) guidance, as well as the CDER/FDA draft guidance. Although neither guideline provides specific recommendations on the conduct of in silico assessments, a recently published survey of industry practice showed that the in silico methods employed by 8 companies are highly similar. More importantly, the survey showed that in silico analysis alone or in conjunction with an expert evaluation provides a high degree of confidence that an impurity that is predicted non-mutagenic will produce a negative result in the Ames assay. Despite the encouraging survey results, there remain specific points of concern which are currently being addressed by a European pharmaceutical industry (EFPIA) working group in collaboration with regulators. The working group intends to provide recommendations on the conduct of (Q)SAR assessments, which, if followed, would be considered sufficiently rigorous methodology. Although many pharmaceutical impurities can be confidently categorized as mutagenic or non-mutagenic based on in silico methodology alone, there is evidence that biological testing is necessary. The Ames assay is considered to be the most practical and appropriate test for the identification of directly DNA reactive substances that warrant TTC control. In most cases the outcome of this assay is sufficient to define and implement appropriate impurity control measures. However, on occasion (e.g. Ames positive degradant that can’t be controlled to TTC limits) additional testing is necessary to further address the biological relevance and risk of Ames positive results. Case studies will be used to illustrate potential scenarios warranting additional testing and factors to take into consideration when deciding what studies to conduct.

### 840 Less Than Lifetime (LTL) Carcinogenic Risk Limits for Mutagenic Impurities during Clinical Development and in Marketed Products.


Mueller et al have provided in 2006 a framework for the application of the threshold of toxicological concern concept to pharmaceuticals with introduction of a “staged TTC”. In this staged approach, acceptable daily intake levels were defined for the clinical development phase of pharmaceuticals, i.e. for up to 1 year intake. In the meantime, this “staged concept” was also introduced into regulatory framework in the existing EU regulatory process on genotoxic impurities. In the ICH M7 group, the principle has been expanded based on the application of Haber’s Rule relating concentration (or dose) and duration of treatment. According to a recent review by Felter et al. (2011), the majority of carcinogens are more potent in rodents if the same cumulative dose is given over a shorter period (e.g. three months) than a longer one (e.g. two years). Although the number of compounds tested in such scenarios is low and it is unclear if these data can be extrapolated to very low exposure scenarios, it was proposed to adjust the acceptable daily exposure to mutagenic carcinogens for shorter durations with an extra uncertainty factor in comparison to longer durations of intake. On the other hand, it was recognized that medication treatment durations are often less than lifetime and depend on many factors (indication, nature of disease being treated and late onset of disease in life). Hence, it appeared unreasonable to propose the lifetime TTC or 1.5 μg/day for mutagenic carcinogens in most marketed pharmaceuticals. In adopting this consideration, it is currently proposed to place clinical trials of longer duration (>1 year) and medical intervention schemes of shorter than 10 years into one category of “up to 10 years” with the assignment of an adjusted TTC of 10μg/day. The presentation will include examples for such clinical trials and pharmaceuticals in use.

### 841 Many Potential Mutagens Used in Pharmaceutical Syntheses Are in Less Potent Classes Than the Carcinogens Used to Derive the TTC, Justifying Higher Levels without Increasing Risk.


The TTC for mutagenic impurities was derived using several worst-case assumptions, and was based on the more potent carcinogens in the public databases. The TTC is needed only when insufficient information exists to estimate safe levels of a mutagen. Most reactive chemicals used in pharmaceutical syntheses are not in the categories of potent carcinogens on which the TTC was based (Delaney, Reg.Tox.
Pharm 49, 107-24, 2007), confirmed here by data from 12 companies and 602 "alerting" structures. Thus, for many synthetic intermediates with structural alerts for mutagenicity, higher daily intakes are appropriate without increased risk. Especially in early stages of pharmaceutical development the risks from exposure to potentially mutagenic impurities are negligible without imposing the default TTC for each. The most common classes are alkylating agents and aromatic amines. For alkyl halides, the relation between the complexity of the structure and carcinogenic potency is well established, (Brito and Muller, in Teasdale, "Genotoxic Impurities", Wiley 2010) and justifies an acceptable daily intake for mono-functional alkyl halides 10 times the default TTC. Aromatic amines have a wide range of carcinogenic potencies, and we lack sufficient knowledge to define the likely potency of each new one. However the structural characteristics of the most potent carcinogens are well defined, so one can rule out that a new aromatic amine is a highly potent one. The "cohort of concern" (COC) was highlighted as so potent that even the default TTC did not provide sufficient control of exposure. The COC comprises aflatoxins and N-nitroso compounds, not usually relevant to pharmaceuticals, but also alkyl structures. The known alkyl azoxy compounds are thought to be mutagenic by forming carboxyls and alkylating DNA. In pharmaceuticals, aromatic azoxy groups are used, which cannot form the alkyl carboxylation, so are not in the highly potent class and need not be excluded from the TTC/ADI approach.

There has been an evolution of risk assessment and regulatory guidance for multipletoxic/carcinogenic impurities in pharmaceuticals. While procedures from other relevant industries have been adapted to pharmaceutical impurities (e.g., food, environmental), there are also unique aspects applied to pharmaceutical process development. Scientific discussions have included application of a probabilistic risk assessment to low level genotoxic substances, impacts of structurally/mechanistic similar impurities versus non-similar impurities, and the risk of synergism/potentiators. The science has shaped regulatory guidance to ensure product quality and patient safety throughout development and in registered use. The European Medicines Agency (EMA) and US Food and Drug Administration (FDA) have developed guidelines (USFDA is in draft) which include control of multiple genotoxic/carcinogenic impurities. The International Conference on Harmonization (ICH) is developing guidance (i.e., ICH M7) on DNA reactive (mutagenic) impurities. According to the ICH M7 concept paper, one of the issues to resolve is multiple impurities. Industry is developing strategies to implement these regulatory guidelines. In conclusion, there are many challenges and issues to resolve, but the advancement of risk assessment for multiple genotoxic/carcinogenic impurities will best occur through science and regulatory guidance.

Reducing sodium intake is a critical public health need for all Americans. In spite of nearly 40 years of voluntary effort to reduce sodium intake, Americans average daily intake of more than 3,400 mg of sodium (approx. 1.5 teaspoons) exceeds the existing maximum intake level 2,300 mg/d (1 teaspoon of salt) established by the 2005 Dietary Guidelines for Americans. High sodium intake puts the population-young and old, male and female, and all ethnic groups at risk for hypertension, and subsequent cardiovascular and/or renal events. These study committees convened by the Institute of Medicine consulted many sources and was provided additional information and insight at its public meeting and through the committee's website. The committees recommendations were in three categories: Primary; Supporting and Interim. Primary Recommended Strategy: A coordinated approach to set mandatory standards for safe levels of sodium in food using existing FDA authorities to modify the Generally Recognized as Safe (GRAS) status of salt and other sodium containing compounds. Supporting strategies: A nationally organized campaign to educate the public about risks, healthy food choices and support for government, industry and consumer effort; Update nutrition labeling; Training for restaurant/food service operations; Purchasing specifications by large food purchasers; Enhanced monitoring; Research. Interim strategies: Food manufacturers and restaurant/food service operations voluntarily accelerate and broaden efforts; The food industry, government, professional and public health partners should work together to promote voluntary collaborations to reduce sodium in foods. The specific recommendations and rationale for each strategy will be discussed along with the actions taken since the release of the report.

The average normal salt (NaCl) intake in the world is about 9 g per day. The result of a meta-analysis of 167 intervention studies randomly allocating participants to a low salt diet or a normal salt diet indicates that the effect of a reduced salt intake of about 6 g on blood pressure is less than 1 mmHg in people with normal blood pressure and about 3 mmHg in people with hypertension. These effects were contrasted by significant increases in variables associated with a poor survival prognosis, such as renal, aldosterone, cholesterol and triglycerides. It is therefore not obvious that sodium reduction leads to health benefits. In agreement with this, population studies generally cannot confirm that people on low sodium diets have less morbidity or mortality than people on normal sodium diets, on the contrary these studies show a trend towards increased all cause mortality among individuals on low sodium diet.

While there is considerable clinical evidence to indicate that substantial reductions in dietary sodium intake may result in single digit reductions in systolic blood pressure for salt-sensitive individuals, further evidence reveals that these same reductions may cause a significant increase in morbidity and mortality across a broad
range of the population. It is generally agreed that most of these negative health outcomes are a consequence of the increase in renin-angiotensin-aldosterone system activity that accompanies reductions in salt consumption. Other negative health outcomes may result from the changes in nutrient consumption patterns that accompany recommended salt reductions. The ideal level of sodium consumption for individuals will result from a rational balance of positive and negative health outcomes. As new research findings continue to emerge and concern over potential unintended consequences of restricting sodium intake grows, it is imperative to ensure that nutrition policies reflect the most current, robust and scientifically sound research.

**848 The Importance of Population-Wide Sodium Reduction As a Means to Prevent Cardiovascular Disease and Stroke.**

D. Arnett. Department of Epidemiology, University of Alabama at Birmingham, Birmingham, AL. Sponsor: M. Soni.

High blood pressure is one of the leading causes of preventable mortality and morbidity, worldwide. National health agencies and professional societies around the world recommend reduction in dietary sodium as a means to lower blood pressure and preventing CVD. The American Heart Association (AHA) recommends limiting daily sodium intake to less than 1,500 mg/d for all Americans. For the estimated one in three who will develop high blood pressure in their lifetime, a high-sodium diet may be to blame. Many consumers do not realize that sodium is ubiquitous in the environment and found in many unsuspecting foods at levels that are often unacceptable. Consequently, consumers continue to purchase foods that are relatively high in sodium and increase their risk of hypertension and related health risks. There is overwhelming scientific evidence that lowering sodium intake improves cardiovascular outcomes. Even a modest reduction in sodium intake is likely to result in substantial health benefits, especially when it is achieved in the general population. The AHA has undertaken an organizational approach to sodium reduction in the United States. The health risks associated with consuming too much sodium in the diet are clear and cannot be ignored.

**849 Unique Challenges in Biologic Drug Development: Separating Mechanism of Action from Mechanism of Toxicity.**

L. Andrews1 and M. Todd2. 1Genzyme, Framingham, MA; 2Pfizer, La Jolla, CA.

A common safety concern of biotherapeutic agents is toxicity associated with mechanism of action (MoA). The term MoA refers to the specific biochemical interaction through which a drug substance produces its pharmacological effect. This workshop will address areas of significant impact to biotherapeutic drug development where there is a known disassociation of MoA and MoT in toxicity studies and/or in translation to the clinic. Most nonclinical toxicity studies are conducted in healthy animals which may not predict the effects of a biotherapeutic when used in the patients with specific diseases. The latest paradigms for using animal disease models where there is a known disassociation of MoA and MoT in toxicity studies can range from additional clinical monitoring to product development termination. The answer to this question has even greater significance for biopharmaceuticals when first in human (FIH) Phase I trials are conducted in subjects with disease for ethical and practical reasons. Thus animal models of disease are being used not only to establish preclinical POC but also to assess exaggerated pharmacology and toxicity in an attempt to improve relevance and extrapolation of results to the intended disease population.

**850 Introduction to the Mechanism of Action/Mechanism of Toxicity Challenges of Biotherapeutics.**


Toxicities associated with biologics are often attributed to the mode of action (MoA). Mechanisms of toxicity (MoT) can be caused by “exaggerated pharmacology” which can occur as a direct effect of the biotherapeutic on the intended cell type or biochemical pathway, or as an indirect effect on unintended cell types or pathways. The extent of the pharmacologic response can be influenced by numerous factors including, but not limited to, biophysical properties of the biotherapeutic agent such as binding potency, effector function and immune stimulatory properties, drug dose and/or duration, tissue distribution of the target, perturbation of signaling networks and disease state. Since the expected pharmacologic mode of action of biotherapeutics is usually understood, adverse effects or toxicities may be predicted using relevant test systems. To this end, specialized pharmacologic endpoints are needed in addition to the conventional nonclinical safety assays for toxicity testing in animals. Important considerations to evaluate MoA/MoT include the use of alternative animal models, safety biomarkers, drug exposure and immunogenicity, impact of drug delivery routes and modalities, novel drug formats and the impact of process changes and engineering on drug safety. Challenges for nonclinical toxicity testing of biotherapeutics include how and when to incorporate “fit-for-purpose” investigational approaches, and importantly how to extrapolate in vitro and in vivo data to the clinical setting.

**851 Use of Animal Models of Disease in Safety Assessments of Biotherapeutics—Is This the Future?**

I. Cavagnaro. Access Bio, Boyce, VA.

A guiding principle in the design of preclinical safety studies is to parallel as closely as possible the clinical conditions of use and exposure. In accordance with this principle, much attention is paid to the dosing regimen with respect to route of administration, duration of treatment and dosing interval. With respect to mirroring the characteristics of the patient population to be exposed normal animals appropriately parallel the typical Phase I population in normal subjects (healthy volunteers). Where toxicologists deviate from the principle of correlation of clinical conditions of exposure is in the evaluation of potential toxicity in patients in the later safety and activity/efficacy clinical trials. The deviation from this principle relates primarily to the physiological state of the clinical populations involved as they are no longer healthy volunteers but rather individuals who either have a specific disease and/or are very ill. Therefore a relevant question for the toxicologist to ask is whether toxicity studies in “normal animals” adequately assess the risks in “sick people”. The answer to this question has even greater significance for biopharmaceuticals when first in human (FIH) Phase I trials are conducted in subjects with disease for ethical and practical reasons. Thus animal models of disease are being used not only to establish preclinical POC but also to assess exaggerated pharmacology and toxicity in an attempt to improve relevance and extrapolation of results to the intended disease population.

**852 Antidrug Antibodies: Interpreting Toxicology Studies and Translating Nonclinical Findings to the Clinic.**

C. Horvath. Genzyme, Framingham, MA.

Monoclonal antibodies and other biotherapeutics are an increasing proportion of drugs being developed by pharmaceutical companies. Unintended effects of biotherapeutics on peripheral blood cells may occur in nonclinical testing or only after

**853 Unexpected Toxicities of Biotherapeutics on Peripheral Blood Cells.**


Monoclonal antibodies and other biotherapeutics are an increasing proportion of drugs being developed by pharmaceutical companies. Unintended effects of biotherapeutics on peripheral blood cells may occur in nonclinical testing or only after...
clinical trials or product launch. Nonclinical studies generally predict clinical het-
matotoxicity for recombinant cytokines and growth factors, but have not been predic-
tive for the majority of biotherapeutics with clinical hematologic liabilities, espe-
cially those that are immune mediated and/or of low incidence. Hematotoxicities
may be species-specific and affect one or multiple blood cell lineages. Mechanistically, hematotoxicity caused by monoclonal antibodies or other biother-
peutics can be directly related to the activity of the test article, or can be indirect
generally due to immune-related events. Direct effects can be due to binding to the
intended target or to an off-target molecule. Recently, several publications have
demonstrated species-specific hematotoxicity due to off-target binding; these ef-
fects have been observed in both rats and NHPs. Indirect hematotoxicity is often
due to autoimmunity, biological cascades, antidrug antibodies, or other immune
system responses. Hematotoxicities due to biological cascades generally involve me-
diators of systemic inflammation, such as cytokines, complement, and immune
complexes, and often are observed as acute post-dosing events. Platelets are particu-
larly susceptible to effects of biotherapeutics, perhaps due to the high expression of
activating FcRs. In vitro assays can be utilized to further investigate and understand
the pathogenesis of unexpected effects of large molecules on blood cells. Characterizing the hematotoxicity (e.g. primary or secondary, relevance to humans,
underlying mechanism, monitorability, etc) is important for risk assessment. Despite
the potential for unexpected hematologic toxicity, the risk-benefit profile of most
biotherapeutics is favorable; hematologic effects are readily monitorable and
managed by dose modification, drug withdrawal and/or therapeutic interven-

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tion.

The 21st-Century shift to more prospective hazard identification and hypothesis
 generation requires greater strategic application of systems biology. QSAR and
archived toxicological data in the form of adverse outcome pathways (AOPs). AOPs
describe the causal linkages among biological responses to chemicals over time. The
complexity of integrating science can be a barrier to progress in terms of the toxic-
ity pathways and networks involved as well as the need to organize knowledge from
many disciplines.

Effectopedia is an open-knowledge aggregation and collaboration tool for delineat-
ing AOPs in an encyclopedic and predictive manner. It includes discrete cause-e-
effect studies and critical reviews that are relevant to toxicology. To achieve human
and machine interpretability, Effectopedia uses an ontology-enhanced, natural lan-
guage interface that offers clarifying questions and specific tags to define the semantic
knowledge while preserving the natural language description of the AOP’s ele-
ments. The use of ontologies also allows Effectopedia contributors to publish their contributions as nanopublications.

Effectopedia serves as a graphical editor to delineate causal linkages at any level of
biological organization and test species. It creates a common organizational space that
(1) helps experts identify gaps in knowledge of causal linkages of biological re-
sponses and (2) acts as a web-based conference room for dialogue and synthesis by
experts with interest in specific AOPs. Effectopedia’s live documents are instantly open for focused discussions and feedback, whilst giving credit to original authors and reviewers. New contributions are immediately distributed to interested parties, keeping all information current and documented. Uncoupling the contribution and review processes also permits organizations to define their own seals of approval and associate them with special interest pathways without slowing down the Wiki-
inspired stream of contributions.

The Chemical Evaluation and Risk Estimation System (CERES) project at FDA Center for Food Safety and Applied Nutrition develops a foundation providing workflows for decision support activities for both pre-market and post-market
safety assessments of food additives, food contact substances, and potential con-
taminants. CERES v1.0 is a data repository and platform for database searching
and computational tools. The content includes chemical structures and properties,
regulatory records, toxicity studies, and other biological screening assays. CERES
is also an institutional knowledgebase where historical regulatory decisions on a given
substance are housed. The system also assists new decision-making processes in a sys-
tematic and consistent manner. In cases where no information is available for a par-
cular substance, CERES provides tools to estimate the toxicity and to assist with
other aspects of safety assessment. Two use cases of the CERES system are described in
this presentation. One is the read-across process within the CERES system to esti-
timate a carcinogenic potential of a new chemical. The analog searching capability
along with the supporting experimental data as well as genetic toxicity and carcino-
genicity models based on mode-of-actions are employed. Various information types
are combined using a weight-of-evidence (WOE) decision theory method. The sec-
ond use case is a post-market analysis of all historical food-contact substances in the
database of the Office of Food Additives using the MOA (mode-of-action)
QSAR models and the CERES toxicology endpoints, carcinogenicity, and
reproductive-developmental endpoints. In addition, a plan to connect CERES to
other predictive methods to enhance the system will be discussed.
Flexible and Transparent Computational Workflows for the Prediction of Target Organ Toxicity.

A. N. Richarz1, S. J. Enochs1, M. Hewitt1, J. C. Madden1, K. Przybylak1, C. Yang2, M. R. Berthold1, T. Meini1, P. Ohl1 and M. T. Cronin1. 1School of Pharmacy and Chemistry, Liverpool John Moores University, Liverpool, United Kingdom; 2Mabion LLC, Columbus, OH; KNIME.com AG, Zurich, Switzerland.

In silico modeling of target organ toxicity has been held back in part by an inability to capture all relevant information into a meaningful reductionist approach. It has also been considered at times too simplistic, using data of often variable quality and seldom allowing the user to assess the relevance to the intended use. The purpose of this study was to develop a novel computational toxicology workflow system, to allow the users greater control and understanding of the target organ toxicity prediction. The workflows were built on the KNIME open-access platform which allows pipelining via a graphical user interface. Various building blocks, known as nodes, were incorporated, to access chemical inventories and/or databases, to produce structures and calculate properties and to report prediction results. The “basic” user sees a web interface, whilst a “trained” user can go behind this to interrogate the nodes and, if required, link to additional data sources or investigate and update the models. The workflow was developed to address in particular the prediction of target organ toxicity of cosmetic ingredients. It comprises an inventory of over 4,400 unique chemical structures (cosmetic ingredients and related substances). The database contains repeat dose toxicity data for over 1,100 compounds including NOEL values. Thus, a user is able to search for similar compounds in the information provided by the workflow: “Pipeline” (levels 1-4) and “Curve fit” (levels 5-8). The pipeline portion from raw data to finalized hit calls and is distributed within two major portions of the ToxCast program with the aim of providing high-quality screening data on thousands of chemicals. The huge library of compounds and wide diversity of high-throughput assays in ToxCast present daunting practical challenges with respect to data processing, analysis, and management with the goal of providing transparent and sustainable data outputs across these massive data sets. To successfully address these goals, the data analysis workflow was designed with 8 levels of processing from raw data to finalized hit calls and is distributed within two major portions of the workflow: “Pipeline” (levels 1-4) and “Curve fit” (levels 5-8). The pipeline portion takes incoming raw data files, performs automated chemical and assay mapping, provides plate/batch effect correction, and assay specific data normalization to supply highly consistent data formats for the curve fitting process. The curve-fitting frameshifts and base substitutions. Later, many strains with the same target sequence but improved antibiotic resistance, cellular permeability, and reporter mechanism were introduced as more sensitive alternatives. However, most existing Quantitative Structure-Activity Relationship (QSAR) models predict either mutations in major strains or overall mutagenicity. We hypothesized that using additional sensitive strains can provide more refined models predicting sequence-specific mutagenicity. Hence, we presented the Chemical Carcinogenesis Research Information System (CCRIS) database (n=6,545) and the CASE Ultra dataset (n=5,438) to create a larger database of 6,967 molecules with fewer missing entries (42% vs 33-39% in original sources). Using mutagenicities reported either in the presence or absence of metabolizing enzymes, we built QSAR models for five sequence-specific mutation sites (hisD3052, hisD3018, hisD6610+hisO1242, HisG46 and hisC3076). These 10 models were compared with the previous strain-specific models in CASE Ultra. Sequence-specific models generally exhibited greater sensitivity (68-88%) and dataset coverage (88-98%) but lower specificity (51-72%). Conversely, previous strain-specific models, built on unbalanced data sets, exhibited high specificity (82-99%) but low sensitivity (17-82%). We have further analyzed the models in terms of statistically significant chemical fragments that were interpreted as structural alerts, highlighting the susceptibility of targeted DNA sequences to certain chemical features. For example, fluorones, due to their planar structures, are good intercalators often associated with frameshifts in hisD3052. The curated data, models, and new structural alerts will be included in future releases of CASE Ultra.


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The representation of structural moieties that carry biological activity information, historically referred to as structural alerts, has a long association with Structure Activity Relationships (SAR). Typically, structural knowledge development is performed only by experts and implemented using SMARTS or proprietary formats. With few exceptions, scientists have generally not been able to query structures with customized substructure rules. Furthermore, there are needs for standardized representations and the ability to encode attributes such as atomic and/or bond properties within structural constructs. Recent efforts to apply chemoinformatics to systems-biology have called for innovation in this SAR area by integrating properties with structure connectivity. To this end, an XML-based representation of chemotypes has been developed to define substructures in which each atom and bond can be annotated with atomic, bond, and/or electronic properties, adhering to a controlled vocabulary, that go beyond the confines of conventional structural classes. These chemotypes have been implemented in a publicly free software application to enable search and data filtering. This chemotype has been developed as part of a project from FDA’s Center for Food Safety and Applied Nutrition to aid in updating FDA structure categories used by toxicology reviewers and to support the mode-of-action modeling (MOA) approach adopted in the Chemical Evaluation and Risk Evaluation System. A series of chemotypes have been developed to incorporate SAR-informed annotations associated with phenotypic effects in hepatotoxicity and developmental morphology, as well as skin irritation and sensitization. These chemotypes enhanced accuracy of categorizations and MOA model predictions. This abstract does not necessarily reflect EPA policies.

In Silico Models of Sequence-Specific Mutagenicity: Exploring Chemical-Mutation Site Associations.

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The Ames Salmonella test for genotoxicity has undergone several transformations since its introduction in 1971. The initial strains, histidine-deficient mutants involving different mutation-probe DNA sequences, were designed to detect specific
and hit-calling process consists of cytotoxicity-point and outlier detection/masking, concentration activity estimation using dose response modeling, systemic data curation and hit-calling process (e.g., fluorescence or non-specific activity). The finalized results are subsequently uploaded into ToxCastDB for data integration and analysis as well as to serve as the primary portal for publication and data release. In addition, the ToxCast workflow has data quality and error identification features quantifying assay performance and quality. The ToxCast data workflow has been systematically developed to efficiently address data management challenges, increase curve-fitting and hit-calling accuracy, and to allow for technology specific customization resulting in repeatable and transparent analyses. This work does not necessarily reflect Agency policy.

863 High-Throughput Electronic Literature Libraries (E-Libraries) to Support Development of Toxicity Prediction Models and Adverse Outcome Pathways (AOPs).

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The elucidation of adverse outcome pathways (AOPs) and mechanistic models of toxicity depends on a thorough understanding of the relationships among protein targets, molecular pathways, cellular processes, adverse outcomes, and exogenous chemicals in a given biological system. Scientists rely on the biomedical literature for a significant portion of this information; however, common tools and resources such as PubMed are designed primarily for article retrieval and may fall short in synthesis of important relationships from the large amounts of available literature. We developed a series of electronic libraries (e-libraries) to provide investigators a way to streamline information retrieval from the biomedical literature and focus the retrieval on key relationships. The foundation for the e-libraries is a broad literature database in the form of a compilation of the MeSH annotations extracted from each MEDLINE record. These annotations are processed and organized so that the relationships among target proteins, biological effects, and chemicals are more easily extracted. For a particular area of interest (e.g., cleft palate, limb development, endocrine disruption) or for a particular set of chemicals (e.g., ToxCast) the literature database is batch-searched for articles containing the corresponding MeSH terms and the co-annotated terms. To produce the e-library, the terms are organized and written to Excel spreadsheets. Finding relevant articles is facilitated by allowing users to use built-in filtering capabilities. A filtering flag for species for instance, will allow the user to see only the rows about a desired species. The e-libraries have been constructed around subject areas as diverse as cleft palate, zebrafish, retinal development, flame retardants, and endocrine disruption. Access to e-libraries has been used to assist all kinds of research from basic exploratory research to development of AOPs and computer simulations. [This abstract does not necessarily reflect EPA policy.]

864 Interactive Web Application (Dashboards) to Integrate Data on Chemical Hazard and Exposure.


The USEPA has developed web-based interfaces (Dashboards) to synthesize multiple sources of data from Aggregated Computational Toxicology Resource (ACToR) into customized information displays. Users may select subsets of chemicals and data types to focus on for the assessment process. The data are organized into custom classes to present all relevant information for a given task. While not all chemicals will have complete information across all classes, each class reports available data including relevant high-throughput assay results, regulatory information, relevant articles from the biomedical literature, exposure data, and in vivo toxicity data. Custom widgets display each type of data according to its nature (e.g. dose-response plots or tables for assay data and summary tables for exposure data). Chemical-specific summary scores that consider all information in each class are presented in a dynamic summary table that highlights areas deserving additional attention. Transparency in decision-making is preserved since Dashboards record and save all user selections and decisions. Expert judgment can be used to override score criteria if necessary, again with such events recorded for transparency purposes. The scores are carried over into a prioritization process where the chemicals are ranked in a weighted-evidence scheme to identify candidates for focused assessment. For example, the Dashboard created to support analysis of potential endocrine disruptors contains over 8000 chemicals with the classes Estrogen, Androgen, Thyroid, Steroidogenesis, Exposure, Occurrence, and Health Effects. The aggregation of relevant data and use of Dashboards to generate custom displays streamlines the toxicity assessment process and allows users to zoom from larger chemical subsets down to very focused chemical-wise views. While the initial Dashboards have been created to support EPA programmatic needs, future versions will include a public-facing web interface. This abstract does not necessarily reflect US EPA policy.

865 Literature Visualization and Navigation to Facilitate Development and Application of Computational Biological Models.

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Physiologically based pharmacokinetic (PBPK) and quantitative structure activity relationship (QSAR) modeling rely heavily on mining of published literature for model development and parameterization, a time and resource intensive process. Because PBPK and QSAR modeling are relatively new disciplines, there is a need for readily accessible and high quality resources to support the increasing model use. Resource needs are broad, ranging from parameter values to model archives and publication archives. A common thread is the need to organize the available model related literature in a fashion that supports rapid access and organization according to data needs and more detailed analysis. We have developed a tree map application, a powerful, visually intuitive literature navigation interface, which exploits the naturally hierarchical nature of information used in biological modeling. Tree maps organize data thematically, and can be navigated by the user choosing a theme of interest, and then drilling down through sub-themes to the level of individual documents (existing models or relevant publications). Subject matter experts developed a taxonomy of themes in biological modeling, and relevant documents were identified in the literature by means of specific search strings. IN-SPiRE Visual Document Analysis was used to identify linguistically significant themes within the data, and to develop and test search strings used by the tree map to identify relevant documents from within the database of biological modeling publications. Documents were then labeled with their relationships within the class structure, loaded into a database, and a tree map user interface generated. With continuing refinement of search strings and development of the tree map application, this work has the potential to yield a powerful, evergreen (automatically updating) search tool specifically designed to aid in biological model development and utilization. An external demo of the tree map can be found at https://nvacademo.pnl.gov/srs/dashboard/epa/

866 Database of Chemicals Assayed for Estrogenic Activity in the US FDA's Endocrine Disrupting Knowledge Base (EDKB).

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Endocrine disruptors (EDs) are chemicals with ability to disrupt endocrine system. EDs have sparked intense international scientific discussion and debate and provoked regulatory action. The FDA's National Center for Toxicological Research (NCTR) developed a publicly available Endocrine Disrupting Knowledge Base (EDKB) (http://goo.gl/Qn0nv) 13 years ago. It has been a widely used resource in the scientific community. Currently, we are expanding the data collection in the EDKB by including the most up-to-date literatures. Since many EDs display estrogenic activities and affect the normal estrogen signaling pathway, we first updated the estrogenic data and developed a database of chemicals assayed for estrogenic activity as a component in EDKB. The database contains over 16,500 activity data of about 7,000 compounds from many public resources. The activity data are categorized based on assay types into binding assay, reporter gene assay and cell proliferation assay. The database also contains protein subtypes and domains used in the assays. Original data were converted into the same units such as logRBA for binding data (RBA=Relative Binding Affinity). Quality control was performed to ensure the data quality. The database was implemented using Instant JChem. A user-friendly interface was designed and developed for rapid navigating and searching the database. Searching multiple fields such as chemical structure, substructure, chemical name, CAS, assay type, assay species, activity range, and references can be conducted using logical operations. To the best of our knowledge, this database is the most comprehensive collection of chemicals assayed for estrogenic activity. It provides the scientific and regulatory communities a free source to search estrogenic activity data for chemicals of interest and to develop predictive models for predicting chemicals for which no assay data are available.

867 The Comparative Toxicogenomics Database.

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The goal of the Comparative Toxicogenomics Database (CTD; http://ctdbase.org) initiative is to inform hypothesis development about mechanisms of chemical actions and the impact of environmental exposures on human health. CTD is a freely

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available tool that provides a foundation of manually curated data describing chemical-gene interactions and chemical-disease and gene-disease relationships. Curated data are integrated with external data sets (e.g., Gene Ontology and pathway data) and tools that allow users to navigate and analyze over 16 million biological relationships relevant to environmental health sciences (EHS). Here we describe CTD functionality and highlight several new analysis and data visualization features, including: a Gene Entrech tool that provides deeper insight into the functional themes and pathways associated with user-defined genes or genes forming the basis of inferred chemical-disease relationships; filtering options that allow users to better customize queries and analyses; and a ‘slim’ list for our disease vocabulary that was implemented with graphical views to provide visual summaries of disease categories associated with chemicals in CTD. We also present information about emerging projects, including the use of curated data for developing chemotypes (a chemical substructure that carries information about biological activity), text-mining workflows for identifying chemical-gene-disease data, novel pathway development tools, and exposure data curation. We continue to expand the depth and functionality of CTD to meet the evolving needs of the EHS research community.


Automated natural language processing (NLP) will make the practice of risk assessment more efficient. In the future, computer algorithms will use NLP to screen the literature for appropriate studies, identify relevant data, and help synthesize mode of action (MOA) evidence across several studies. However, before we can use the data from these searches, ontologies must be built to facilitate efficient data storage and sharing using existing standards. We developed the Mode of Action Ontology (MOAO) to support automated NLP discovery, integration, and management of mode of action knowledge. MOAO is a semantic model that 1) describes MOA knowledge, 2) easily combines data from multiple sources, 3) makes MOA knowledge more transparent and easily available to the public via semantic web interfaces, and 4) will allow computers to discover information and relationships via logical inference. MOAO is the first ontology we are aware of that describes mode of action information for toxicological purposes. MOAO was conceptualized as an application ontology, borrowing terms from existing high-quality ontologies produced for various biological sub-specialties. This allows us to maximize integration with outside data sources. In this way, MOAO can be a model for how other sub-domains in toxicology can reap the benefits of ontologies without the overhead of constructing one completely from scratch. We leverage the work by the Open Biological and Biomedical Ontologies (OBO) Foundry and the Basic Formal Ontology (BFO) to facilitate the integration of our ontology with others. The reference ontologies which MOAO uses include the Gene Ontology (GO), CHEBI (Chemical Entities of Biological Interest), the Uber Anatomy Ontology (UBERON), the NCBI Taxonomy, and the NCIT Thesaurus. The views expressed in this abstract are those of the authors and do not necessarily represent the views or policies of the U.S. Environmental Protection Agency.

Development of a COSMOS Database to Support In Silico Modelling for Cosmetics Ingredients and Related Chemicals.
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The Seventh Amendment of the EC’s Cosmetics Directive foresees a deadline in 2013 for the replacement of animal testing of cosmetic products for repeated dose/ repetitive toxicity and toxicokinetics. To this end, the COSMOS project has been initiated to fill the gap in knowledge and technology via in silico toxicology. The management and sharing of chemical, biological and toxicity data play a central role in any database designed for in silico toxicology: an overview of toxicity data sources and studies; the design of the data model and the data entry tool for the database; and the definition of a data curation strategy. The new COSMOSDB provides a publicly available web-based searching/ retrieval system and also serves for sharing resources, models and supporting workflow developments. The database is “compound-centred” and represents both EU COSING and US PCPC inventories for cosmetics ingredients. For biological data, the database supports repeated dose toxicity, metabolism information and dermal absorption data. The data have been collected, curated, quality-controlled, stored and managed in a flexible and sustainable manner to support predictive modelling tasks. An in vitro toxicity data set has also been reflected in building the data entry system. Based on the review of existing approaches on good practice to assess quality entries, the reliability of the toxicity data is supported by all available data from multiple sources. Additional functionalities for data aggregation within a data governance framework are described. The COSMOS project is funded by European Community’s 7th Framework Program (FP7/2007-2013) and by Cosmetics Europe.

The aim of the €50 million SEURAT-1 (Safety Evaluation Ultimately Replacing Animal Testing-1) research cluster, sponsored by the European Union and Cosmetics Europe, is to generate a proof-of-concept replacement for repeated dose systemic toxicity testing on animals. The SEURAT-1 strategy is to adopt a mode-of-action framework to describe repeated dose toxicity, combining in vitro and in silico methods in an intelligent manner to derive predictions of in vivo toxicity responses. ToxBank is the cross-cluster infrastructure project whose activities include the development of a data warehouse, the selection of reference compounds for use across the cluster to support the mode-of-action framework, a physical compounds repository, and a reference resource for biomaterials. The ToxBank data warehouse provides a web-accessible shared repository for protocols and experimental results supporting SEURAT-1 research objectives such as toxicity pathway elucidation and biomarker discovery. Information in ToxBank is uploaded from the research activities of the cluster partners and relevant information on the reference compounds are drawn from public databases. In this poster we describe the data warehouse and implementation based on open standards, the selection of reference compounds, and an analysis linking the reference compounds to public omics data.

Software for Integration of Experimental, Chemical, and Environmental Data.
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The Oregon State University Superfund Research Program evaluates the potential health effects of polycyclic aromatic hydrocarbons (PAHs). More than 100 parent PAH compounds, and an unknown number of PAH derivatives and metabolites have been identified. PAHs raise concern at over 800 Superfund Sites and serve as the risk driver for remediation at 20% of all Superfund sediment sites. Assessing the health effects of PAHs is a major challenge because environmental exposures to these chemicals are from complex mixtures of PAHs and other toxicants. Integration of exposure data with biological activity and pharmacokinetics is essential to evaluate the effects presented by these mixtures on human health. Increasingly researchers are generating data streams of high content data from analytical, high-throughput, and ‘omic’ studies, which make direct applications from these technologies out of reach for clinicians and regulators. However, more accurate and precise thresholds for risk assessments and prediction will ultimately arise from leveraging the integration of data from across disciplines. Therefore, the principle challenge for mixture assessments, espoused by PAHs, is the efficient integration of large and diverse data sets. To this end we’ve extended PNNL’s Experimental Data Management System (EDMS) to create a seamless data workflow among researchers and analysts. EDMS has been extended to track and manage relationships among any and all data points across a project. This allows us to integrate chemical and biological data for environmental extracts (LIMS and EDMS) with analogous data on pure chemical compounds. EDMS provides a query mechanism that will reach through all the relationships of data from chemical compounds to the analytics and statistics run on the experimental data. Query results, visualizations, and data processing results are all also stored together with and related to the experimental information. This reduces data replication and supports a streamlining of analysis, collaboration and experimental design allowing researchers to take advantage of all the data generated.
Drug development scientists and regulatory reviewers alike have a need to understand toxicity patterns of a drug candidate across animal species prior to human trials. Effective exploitation of collected study data has previously been impeded by differences in nomenclature, terminologies and units across sponsor and CRO data. Moreover, computational techniques to analyze large volumes of disparate quantitative and qualitative study data, and their respective endpoints, have not been routinely available for use by practicing toxicologists and reviewers. To address these common challenges for the FDA and the Industry, we have developed new repository techniques to prepare and hold in vivo harmonized datasets (including standardized formats like SEND, the Standard for Exchange of Nonclinical Data), and extracted metadata, as a first step towards advanced analytics and study reviews. Predictive, comparative and cluster analysis techniques have been performed on large study repository datasets to reveal similarities and differences in dose-response patterns across species in terms of their magnitude, shape and trends. Finally, signal processing techniques such as Z-transform box plots have been adapted to present trends in ways that scientists and regulatory reviewers can easily use to interpret potential toxicity patterns and their inter-relationships. Collectively, new computational tools are available to better exploit legacy and ongoing data that are already being collected, provide better decision support for species selection and protocol design, and advance our understanding of how dose-response patterns translate across species for the same compound and to detect other compounds that share similar responses. Implications of applying such computational methods to improve safety prediction across species, and eventually into humans, will be discussed particularly in light of the FDA’s mandate to require standardized e-Data submissions, including SEND datasets, as a way to further regulatory science.

**Purpose:** Purpose of this study is to develop and apply a novel technique of combining the predictions of the CASE Ultra expert system models and a new quantitative read across modeling methodology to obtain better genotoxicity predictions as opposed to using just one technique. The hybrid methodology successfully provides the interpretability of toxicity alerts and reduces false positives, false negatives while increasing the test coverage. This in silico tool can be successfully used for assessment of potential toxicity or beneficial therapeutic effects of new drug candidates, impurities and metabolites.

**Method and Data:** CASE Ultra is a computer program that automatically extracts structure-activity knowledge from chemical databases and applies the tools for predicting activity in test chemicals. It also gives a very clear explanation of its predictions so that they are easily interpretable by chemists and toxicologists. Recently we have developed a novel read across modeling technique for quantitative prediction of toxicity that uses the full neighborhood profiles of chemicals. In case of Salmonella mutagenicity predictions, the read across prediction was applied in a systematic way on top of the predictions from the CASE Ultra mutagenicity models, and significant improvements in the sensitivity, specificity and coverage were achieved. The details of the methodology and results of prediction for a large external test set are presented.

**Results of the Study:** The improvements in the predictive performance as a result of the hybrid technique is demonstrated for Salmonella mutagenicity prediction in an external test set. Improvements in the prediction sensitivity, specificity and coverage were in the order of 3-5%, 2-5% and 5% respectively.

**Conclusion:** The hybrid methodology successfully produces predictions that are comparable to the predictions from CASE Ultra. Future work will focus on improving the interpretability of the hybrid methodology to support regulatory decision making.
We propose developing a modular, cloud-ready, informatics-based system to synthesize multiple data sources into overall human health assessments of chemicals. This system would seamlessly integrate and document the overall workflow from literature search and review, data extraction, and evidence synthesis, to dose-response analysis and uncertainty characterization. Crucial benefits of such a system include improved data integrity, greater transparency, standardization of data presentation, and increased consistency. By including both a web-based workspace for assessment teams, and complementary web-based portal for reviewers and stakeholders, all interested parties would have dynamic access to completed and ongoing assessments. The modular approach will also facilitate rapid prototyping, testing, review, and incorporation of methodological improvements. Here we present a prototype module for benchmark dose (BMD) modeling used to develop points-of-departure, from which toxicity values are derived. Previously-developed BMDS Wizard and DRAGON Excel-based programs were used to develop a web-based tool where assessment teams can view/upload/enter dose-response data sets into the module, perform BMD modeling, and export results. Example summary views and plots are available online, or can be converted to report format. In addition, multiple nested views of the data and analyses enable interested users to rapidly "dive into the details." We conclude that given new data streams, diverse user needs, and multiple stakeholder interests, assuring the utility, integrity, and objectivity of human health assessments will be greatly facilitated by a modular, upgradeable, informatics-based system for their development, review, and dissemination. Disclaimer: The views expressed in this abstract are those of the authors and do not necessarily represent the views or policies of the US or California EPA.

Cellesis: A Novel System Biology Toxicology Tool

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Understanding and predicting adverse drug reactions (ADRs) are crucial goals in toxicology, since unrecognized toxicity is the most frequent reason for drug withdrawal. We have developed a novel technology based on dynamical image analysis of intracellular organanelles in live cells. Mitostream®, the dynamic measure of the mitochondrial behavior, is a novel technology predictive of the severity and incidence of ADRs. It is a powerful HCA to help identify drug-induced toxicity, differentiate risk between compounds and evaluate the potential for rare but severe adverse events. Mitostream® decodes the mitochondrial behavior, quantifying motility, morphology, permeability and network organization with multiple endpoints associated to qualitative descriptors to model their interrelations. We developed the Cellesis algorithm to identify classifiers and predictors able to fuse the matrix of recorded multi-modal markers into a unique statistical toxicity index. An original set of 27 molecules was studied to classify drugs on a toxicity scale. The algorithm classified properly 83% of the drugs with 94% specificity and 75% sensitivity. We extended the set to include more non-toxic compounds and compounds with rare but severe toxicity as well as compounds that did not target the mitochondria. The second training set included 47 compounds and yielded 85% proper classification, a specificity of 91% and a sensitivity of 92%. Independent industrial validation was performed through the Consortium for Technology Evaluation at the DSECS, testing 10 new compounds. Chosen by our industrial partners these compounds classified properly with similar performances, showing a good association between Cellesis-predicted ADRs and the lowest tolerable dose seen in the clinic. A set of NSAID was further studied. Results showed 86 % proper classification through Cellesis-predicted ADRs. Specificity was 92%, sensitivity was reduced to 85%, some apparent false positive had a hollow threshold reported clinical ADRs despite having been withdrawn post-marketing due to idiosyncratic hepatoxicity.

Development of a Systems Computational Model to Investigate Early Biological Events in Hepatic Activation of Constitutive Androstane Receptor (CAR) by Phenobarbital.


Activation of the nuclear receptor CAR (constitutive active/androstane receptor) is implicated in the control several key biological events such as metabolic pathways. Here, we combined data from literature with information obtained from in vitro assays in the US EPA ToxCast database, and in vivo outcomes in the US EPA ToxRef database to link CAR activation to a proposed adverse outcome pathway (AOP) of hepatic tumors. Key in vivo pathological events for this proposed hepatic tumor AOP are similar to those of other non-genotoxic receptor-mediated chemi- cals (i.e., hypertrophy, cellular proliferation, preneoplastic foci, and neoplasia), highlighting the importance of upstream biological mechanisms mediating the early proliferative signals of CAR activation. Using phenobarbital (PB), a known rodent tumor promoter and CAR activator, we developed a computational model for the early initiating events of CAR activation leading to its nuclear translocation. Based on literature evidence, the computational model describes the cascade of bi- ological events leading to CAR nuclear translocation. These events are initiated by PB activation of Adenyl Cyclase (AC) leading to an increase in the rate of ATP transformation to cyclic AMP. In turn, cyclic AMP increases intercellular calcium levels which inhibit the activity of AC via a negative feedback loop mechanism. However, the resulting transient increase in cyclic AMP levels activates protein.
Mechanistic Modeling Illuminates the Most Important Unknowns in Bile Acid Mediated DILI.

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BSEP inhibition and consequent bile acid (BA) buildup has been proposed to be an important event in drug induced liver injury (DILI). There are many gaps in the knowledge of BA homeostasis and its disruption by transporter inhibitors. Modeling can help us understand which of these knowledge gaps should be filled first in order to provide maximum understanding. We have constructed a model of BA homeostasis and toxicity within DILysim1, a mechanistic model of DILI that includes bile acid flux into and out of hepatocytes, the role of the farnesoid X receptor (FXR) in the regulation of BA transport and synthesis, and gallbladder BA release and recirculation. We first analyzed our system's behavior in the presence of a simulated BSEP inhibitor similar to glibenclamide. We predict a 41% increase in total hepatocyte content of amide conjugated chenodeoxycholic acid (CDCA). However, the simulated BSEP inhibition resulted in no change in either unconjugated CDCA or the amide-conjugated lithocholic acid (LCA), and the hepatocyte content of unconjugated LCA decreased by 15%. Next we performed a sensitivity analysis on our system's response to BSEP inhibition in order to determine which parameter has the greatest effect on hepatocyte content of BAs and hence BSEP inhibitor-mediated hepatotoxicity. We found that the most important unknown in the system is the intrahepatic BA concentration that triggers FXR-mediated regulation of BA transporters, while other parameters such as the affinity constants for transport of CDCA and LCA and ileal LCA transport efficiency were less important. We conclude that the highest priorities for wet lab research to inform our model and enhance our understanding of BSEP inhibitor mediated DILI are: 1) quantifying the relationship between intrahepatic concentrations of individual bile acids and FXR activation, and 2) quantifying the link between intracellular CDCA and the hepatocyte culture model built for TCDD can predict the toxicities or downstream events for other AhR agonist by changing the parameter associated to AhR binding affinity only. This evaluation provides a model example of a quantitative in vitro to in vivo extrapolation for nuclear receptors and informs on how to better use in vitro data for risk assessment. This abstract does not necessarily reflect the policies or views of NIH.

Quantitative Extrapolation of In Vitro Data to Risk Management Decisions: Using Hepatic Ah Receptor and Its Agonists As an Example.

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A critical question in human risk assessment is how best to incorporate in vitro data. In a quantitative in vitro to in vivo extrapolation, one assumption is that media concentration equals blood concentration. Here, we evaluate this assumption by applying physiologically based pharmacokinetic (PBPK) models to predict cellular responses in primary rat hepatocytes exposed to an aryl hydrocarbon receptor (AhR) agonist. PBPK models for AhR agonists (e.g., TCDD, PaCDE, PCB126) were developed based on NTP two-year rat carcinogenicity bioassay data. The model predicts tissue concentrations and CYP1A1 induction well and demonstrates that ligand binding affinity is an important parameter for AhR-mediated enzyme induction and distribution of these chemicals. Using this model, we predicted the dose response for Cyp1a1 induction in whole rat liver at 48 h after an oral dose of TCDD. The predicted results were compared to literature data on rat primary hepatocyte treated with TCDD for 48 hours. The EC50 obtained for Cyp1a1 induction in vitro is about 100 fold less than that predicted in vivo (i.e., the primary cell culture system is about 100 times more sensitive). To identify which factors might contribute to this difference, we built a primary hepatocyte culture model, based on rat primary hepatocyte data from the literature and incorporated the same parameters used in the in vivo PBPK model of AhR agonists. Based on differences in parameter values between the two models, we identified critical factors determining the difference in sensitivity between in vitro and in vivo (e.g., TCDD availability). The hepatocyte culture model built for TCDD can predict the toxicities or downstream events for other AhR agonist by changing the parameter associated to AhR binding affinity only. This evaluation provides a model example of a quantitative in vitro to in vivo extrapolation for nuclear receptors and informs on how to better use in vitro data for risk assessment. This abstract does not necessarily reflect the policies or views of NIH.

Predicting Plasma Profiles following Oral Dosing for Drug Liver Transporter Substrates Using Physiologically-Based Pharmacokinetic Modeling.

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A novel prediction approach for OATP substrates was developed using in vitro sandwich cultured hepatocyte (SCHH) data as input into a physiologically based pharmacokinetic (PBPK) model and was validated for intravenous (iv) dosing (Jones et al., 2012). The aim of our work was to extend this to the prediction of oral profiles. Data for plasma concentrations following oral doses were obtained from literature. Simple descriptors of oral absorption (fa, ka) were predicted using a physiologically based gastrointestinal (GI) tract model, which were then used in the published transporter PBPK model with a single compartment for oral absorption. The seven literature compounds were bosentan, cerivastatin, fluvatat, pravastatin, repaglinide, rosuvastatin, and valsartan. In vitro permeability and solubility under fed and fasted conditions were measured as inputs to the GI model. We tested whether the oral absorption parameters were reasonable by combining them with the transporter parameters obtained from fitting the iv data plasma. Characterizing plasma pharmacokinetics using area under the curve to infinity (AUC(0-inf)), maximum concentrations (Cmax), and time to maximum concentration (Tmax), the fold error for the seven compounds was 1.6, 2.3, and 1.9 respectively, indicating the oral absorption parameters were reasonable. Forward predictions for the seven compounds based on SCHH data and the oral absorption parameters provided a test of both aspects of predicting oral plasma time course. The average fold errors for the seven compounds increased somewhat to 2.5, 3.3, and 2.4, respectively. This method was then applied to oral plasma data for four in-house compounds resulting in reasonable predictions for three of the four. An initial prediction method has been developed using in vitro and in silico inputs to estimate plasma profiles and pharmacokinetic parameters for iv and oral dosing of anionic drugs that are substrates for liver transporters.
A Mechanistic Approach to Understand Idiosyncratic Toxicity of Perhexiline and Sulindac.


Idiosyncratic DILI is one of the most common causes of safety-related post-marketing drug withdrawals. We have developed a dynamic systems based integrative approach that aids in its understanding by combining a mathematical model along with a panel of in vitro enzyme and transporter assays. We tested perhexiline and sulindac in this assay system and fed the assay results into a mathematical model of liver biology. Our simulations identified the key mechanism behind the steatohepatitis caused by perhexiline. They predict perhexiline induced alterations in mitochondrial function, i.e., inhibition of oxidative phosphorylation, generation of oxidative stress along with inhibition of mitochondrial beta-oxidation leads to steatosis. An interesting insight gained in this process was the fact that whole cell measurements such as ATP or GSH alone can provide an erroneous understanding of the impact of a compound in vivo. On the other hand measuring various enzymatic functions enabled us to quantify the impact of ROS in vivo.

In the case of Sulindac, intracellular accumulation of GSH in HepG2 cell line indicated impairment of transport system. Applying this impairment in the model, our simulations predict bile and bilirubin accumulation indicating the potential for idiosyncratic cholestasis. Thus our system enables us to understand mechanistic insights into the effect of leads and make predictions on their effects in vivo without necessitating the use of whole animal studies.

An Integrated Structure-Based Systems Approach to Predict Hepatotoxicity.

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We have developed a systems biology model for hepatotoxicity prediction. The input for the model is a set of in vitro assays that capture the effects of a chemical compound on selected proteins. The model takes these assay results as input, simulates their likely effects on selected pathways in a liver cell and outputs the toxicity effects in vivo. Though this approach has good predictive value and provides mechanistic insights into the toxicity response, the time consuming and expensive assays are a bottleneck for using the model in a high-throughput manner. Using structure based approaches, we have virtualized the assays to use this model to rapidly evaluate hepatotoxicity of molecules. The binding affinity of a molecule to each of the proteins is estimated using structure prediction, flexible and induced docking and pharmacophore approaches and classified as low, intermediate or high. This forms the input to the systems model that simulates the effect of the compound on selected proteins. We found that this method is able to flag the key toxic mechanisms of each of these molecules and is able to predict synergistic toxic effects. We believe that this approach can be used as screening tool to rapidly and accurately assess mechanistic toxic effects and has utility during lead discovery and optimization.

Application of an In Silico Approach to Predict Intrinsic In Vitro Cytotoxicity for Compounds in Primary Human Hepatocytes during Preclinical Development.


Drug-induced liver injury is a major cause of compound attrition in preclinical and clinical development as often, intrinsic hepatotoxicity can be related to chemical structure. Screening in primary human hepatocytes has been routinely used to assess toxicity in the preclinical setting to allow for assessment of metabolic contribution. However, the knowledge gained from this screening has not been fully utilized to drive medicinal chemistry design before compound synthesis. Herein, we present work in progress in the development of a computational model that is used to predict the outcome of hepatocyte screening for a set of preclinical compounds with significant accuracy. Cytotoxicity was determined by treating cryoplateable human hepatocytes with individual compounds and evaluating cellular ATP levels. In vitro cytotoxicity was classified based on IC50 values into three buckets: high (≥50 μM), moderate toxicity (10-100 μM) and low toxicity (<10 μM). Using a machine learning approach, a large set of structural and non-structural descriptors were evaluated for their usefulness to classify cytotoxicity on a training set of 72 compounds. Three calculated physicochemical properties emerged as the most predictive descriptors (most-basic-pKa, plasma protein binding % and logP), and were used to construct a decision tree model. Subsequently, predictions were made for 102 compounds prospectively. The positive predictive values were 77% & 65%, and negative predictive values were 72% & 85% for high (H) and low (L) toxicity groups, respectively. Interestingly, addition of structural descriptors did not improve the accuracy of prediction, suggesting that the intrinsic toxicity of those compounds were not specific to their structures. We show that this model proved useful in reducing compound attrition for an ongoing project. Furthermore, the physicochemical property space that this work has implicated as being associated with toxicity may also provide clues toward understanding the underlying mechanism(s).

In Vitro In Vivo Correlation: Apparent or Ambiguous? Analysis of Pfizer Compounds in US EPA’s ToxCast Chemicals-Assay Space.


In 2009, Pfizer collaborated with EPA in their ToxCast initiative to identify predictive bioactivity signatures for toxic compounds using ‘dead’ pharmaceuticals. A total of 52 Pfizer compounds with preclinical data were profiled in multiple assay platforms measuring bioactivity against diverse assay endpoints. The first part of this work was focused on the analysis of Pfizer compounds in ToxCast chemicals and pharmacological space. These include a) comparison of bioactivity profile of Pfizer compounds with other pharma and EPA compounds; b) analysis of compounds sharing similar chemical and pharmacological space as of Pfizer compounds; and c) ability of assay platforms to reproduce the primary mode of action of Pfizer compounds. Analysis revealed that compound from EPA set seemed to have different pharmacological and chemistry space than pharma compounds. It also highlighted assay platforms and sensitive endpoints that are more relevant to understand the off-target activities of compounds. The second part involves correlation of in vitro ToxCast assay signals with in vivo findings related to liver injury, observed either in preclinical or clinical studies. The analysis pinpointed diverse cytotoxicity endpoints within ToxCast as markers of compound potential to cause preclinical or clinical liver injury.

Improved In Silico Methods to Group Liver Toxics Related to Adverse Outcome Pathways.


Prediction of repeat dose toxicity by in silico methods remains an elusive goal. Current quantitative structure-activity relationship (QSAR) approaches have acknowledged limitations. A more pragmatic approach is to identify relevant molecular initiating events associated with an adverse effect – integrated into the Adverse Outcome Pathway (AOP) concept - and attempt to group chemicals accordingly. Such an approach will have a greater probability of success if targeted at organ level toxicity. The purpose of this study was to develop novel structural alerts to enable to the grouping of potential liver toxicants. Specifically, the utility of current approaches for grouping according to protein reactivity (relating to reactive hepatotoxicity) was investigated along with novel methods to group chemicals and extract usable information from the grouping, placing in the context of relevant AOPs. The objectives therefore, were to: a) provide a better predictive model for liver toxicity, but to improve structural alerts to group compounds to perform read-across for liver toxicity. Data for over 900 compounds classified as either being toxic or non-toxic to the liver were assessed in this study. Attempts to group compounds on the basis of protein reactivity (the OECD Profiler in the OECD QSAR Toolbox) had poor predictivity, only identifying approximately 34% of liver toxicants, whilst nearly 25% of non-toxicants were found to contain a reactive structural alert. Therefore, the liver toxicants were grouped according to structural similarity using the ToxMatch software. Over 80 categories were found by this automated method from which 16 robust categories were manually curated, many of which were associated to mitochondrial toxicity AOPs. The suite of reactive and non-reactive MIEs, defined by structural alerts and chemotypes, provides a basis for grouping potential liver toxicants. Supported by the EU FP7 COSMOS and eTox Projects.
Construction of a Human Hepatotoxicity Database for QSAR Modeling.


Drug-induced liver injury (DILI) represents the second most common reason for pre-market attrition of drug candidates and post-market withdrawal of approved products, prompting the need for better tools to predict these serious effects. Quantitative structure-activity relationship (QSAR) modeling uses computational algorithms in conjunction with large chemical data sets to identify correlations between molecular structural features and biological activity, and lends itself well to the task of improving the prediction of drug safety by exploiting FDA's vast institutional knowledge of DILI and other adverse event (AE) data. This report describes the development of a new method for creating a DILI QSAR training set based on FDA's Adverse Event Reporting System data. The method utilizes the Multi-item Gamma Poisson Shrinker (MGPS) algorithm, the gold standard used by FDA CDER's Office of Surveillance and Epidemiology for AE signal identification, to generate Empirical Bayesian Geometric Mean (EBGM) scores for all drugs in the database binned by MedDRA preferred term. A set of 38 preferred terms (PTs) describing serious hepatic events, such as hepatic failure, hepatocellular injury, and hepatic cirrhosis, was selected based on related physiology and their adverse event counts were pooled to provide a more robust dataset for scoring purposes. An independently obtained verification set of 107 known hepatotoxics with evidence ranging from clinical manifestations (n=93) to market withdrawals (n=14), was used as a calibration set to determine whether the standard EBGM cutoff of 2.0 was appropriate for partitioning the training set. While 92% of the withdrawn drugs were correctly classified, only 50% of the positives with clinical manifestations were correctly classified, emphasizing the need for additional parameters and/or PTs to supplement the AE data used in the training set. This method created a set of over 2000 unique molecular entities that were structurally suitable for QSAR modeling.

Evaluation of Perchlorate Intake on Maternal and Fetal Serum T4 Levels Using a BBDR Model.

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Perchlorate (ClO4- is both a naturally occurring and man-made chemical widely distributed in the environment with low levels detected in food and drinking water. Disturbances in the maternal hypothalamic-pituitary-thyroid (HPT) axis leading to hypothyroxinemia and hypothyroidism have been shown to cause negative effects on the neurodevelopment of the fetus. Downstream perturbations of maternal and fetal HPT axes following ClO4- competitive inhibition of sodium iodide symporter-mediated thyroid iodide (I-) uptake have not been evaluated quantitatively. In order to quantitate effects of dietary I- intake, ClO4- exposure and their interactions on maternal and fetal HPT axis, a biologically based dose response (BBDR) model for HPT axes in the pregnant woman and fetus was developed. The BBDR model includes sub-models for I-, ClO4-, thyroxine (T4) and triiodothyronine (T3). The model was successfully calibrated for euthyroid, marginal iodide deficiency and ClO4- exposure. Serum thyroid hormone changes were predicted for diet a Y- intake ranging from 75 to 250 μg/day and for ClO4- exposures of 0.01 to 1000 μg/kg/day. Model simulations suggest that ClO4- at environmentally relevant ranges of exposure (~0.1 μg/kg/day) does not result in significant decreases in maternal and fetal free serum T4 concentrations for maternal I- intake of 75 to 250 μg/day and for ClO4- exposures of 0.01 to 1000 μg/kg/day. For a daily I- intake of 200 μg/day, the daily dose of ClO4- required to reduce serum free T4 (fT4) levels from representative euthyroid region to a hypothyroxinemic state was estimated to be about 50-fold greater (32 μg/kg/day) than the current reference dose (RJD, 0.7 μg/kg/day). As I- intakes were lowered (150, 100 and 75 μg/kg/day), ClO4- doses required for similar reductions in fT4 levels were reduced to 28, 16, and 4 μg/kg/day, respectively. This BBDR-HPT axis model for pregnancy provides a novel tool for public health assessment for endocrine active chemicals found in food and the environment. Funded by the Air Force Center For Engineering & the Environment through 711 HPW/RHDJ, WPAFB, OH.

A Multiscale Dose-Response Model of AhR Toxicity Pathway Activation in the Human Liver.

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We have developed a spatial, multicellular agent-based computational model of the human liver lobule to study the contribution to hepatotoxic injury from multiple levels of biological organization. This multi-scale "virtual tissue" model combines molecular circuits in individual hepatocytes with cell-cell interactions and blood- vessel uptake of toxicants through hepatic sinusoids, and dosimetry estimated by whole-body PBPK models, to enable quantitative, mechanistic prediction of dose-response. We use the activation of the aryl hydrocarbon receptor (AhR) toxicity pathway in hepatocytes exposed to 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) and other dioxin-like compounds (DLCs) as a case study for data integration and multiscale model development for mechanistic dose-response prediction. By explicitly accounting for basal gradients in AhR expression across the lobule and stochastic uptake of TCDD molecules by individual hepatocytes, we were able to reproduce experimentally observed zonal heterogeneity in TCDD accumulation and cytochrome P450 1A1 expression. A detailed implementation of the AhR signaling and transcriptional cascade in each hepatocyte in this multiscale model enabled us to characterize differential dose response for various AhR-mediated endpoints at the tissue level, including cytoxicity and cell proliferation.

Virtual Liver: Mapping Pathways to Liver Injury by Linking In Vitro Assays to In Vivo Outcomes.


The EPA ToxCast project uses high-throughput screening (HTS) to evaluate in vitro bioactivities of chemicals. Here we are attempting to bridge HTS data with histopathological effects in ToxReD using a data-driven computational approach. Starting with 2,200 guideline studies for 763 chemicals stored in ToxRefDB, we defined 8 broad categories of hepatic histological effects consisting of hypertrophy, cytotoxicity, inflammation, regeneration, steatosis, fibrosis, preneoplasia, and neoplasia. A subset of 560/763 chemicals were screened in 500 ToxCast assays. Univariate statistical analysis found 91 assays in which chemicals had a significantly (p<0.05) greater potency for hepatic effects. We identify mechanistic relationships between the molecular and cellular targets of these assays from literature and use an information-theoretic criterion to calculate the most plausible sequence of key events for each outcome. The results are summarized as a dependency network including molecular initiating events (e.g. AhR, PXR, and PPAR), signal transduction (PI3K/AKT, p53, NfkB, JNK), cellular alterations (oxidative stress, mitochondrial dysfunction, cytoskeletal changes, cell cycle arrest and apoptosis), and histological effects. The analysis also suggests markers of hepatic lesion severity: chemicals that cause hypertrophy without injury have a 3-fold potency (p<0.05) for stress kinase activation; for hypertrophy and injury a 2-fold greater potency for G2/M arrest and p53 activation; and for steatosis a 2-fold greater potency for mitochondrial dysfunction. This work should inform current efforts to translate HTS data from a molecular scale to clinically-relevant health effects. This work was reviewed by EPA and approved for publication but does not necessarily reflect official agency policy.


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Polycyclic aromatic hydrocarbons (PAHs) are ubiquitous, often carcinogenic environmental contaminants generated as by-products of combustion processes. PAHs are found at almost half of Superfund sites, and are drivers for remediation in a significant percentage of those sites. Dibenzo[def,p]chrysen (DBC) is a high molecular weight and highly potent transplacental carcinogenic PAH. We developed the first species- and life stage-specific physiologically based pharmacokinetic (PBPK) models for high molecular weight PAHs that can be used to predict target tissue doses of PAH metabolites important to their carcinogenic mode of action. In rodents, we measured metabolic enzyme activity, rates of metabolism, and pharmacokinetics for critical life stages such as pregnancy and development. Pregnant animals had elevated tissue concentrations of DBC and its metabolites (2 – 4 fold increase in CMAX), and suppressed rates of clearance compared to naïve mice. Using activity-based protein profiling (ABPP), a unique proteomic approach for characterizing functional proteins in complex biological systems, to measure and compare enzyme activity, we found that multiple enzymes important for PAH metabolism have significantly altered activity during pregnancy and development in rodents and humans. Incorporating changes in anatomy and physiology during pregnancy as well as measured reductions in P450 enzyme activity accounted for
observed pharmacokinetic differences in mice. We have extrapolated our PBPK models to the human dosimetry on the basis of human anatomy and physiology, and chemical specific human in vitro metabolic rates. In order to evaluate the extrapolated human model, a controlled human pharmacokinetic study of DBC at environmentally relevant exposure levels (30 ng bolus exposure) has been performed by our collaborators. Initial simulations of DBC concentrations in blood are remarkably similar to those observed in preliminary data from human volunteers.


The increased sensitivity of post-natal day (PND) 21 (weaning) rats to acute high-dose neurotoxicity of deltamethrin (DLM) poses the question of whether children will be more sensitive than adults both of whom are exposed to a much lower dose than rats. Age-related pharmacokinetic (PK) differences are hypothesized to be responsible for the observed differences in rats. We developed a PBPK model for human children based on rat PK and rat and human in vitro data. Our model incorporated species and age-dependent differences in metabolism of DLM as well as physiological changes during growth. The predicted age-dependent changes in brain Cmax correlated well with the maturation of metabolic capacity suggesting the lower metabolic clearance as a mechanism for the increased sensitivity in neonatal rats. The predicted Cmax in brain of PND 21 pups after a single oral dose of 4 mg/kg was 50% higher compared to PND 70 rats. The human model was developed similarly by incorporating maturation physiology and developmental changes in DLM metabolizing enzymes. Unlike the rat, the predicted brain Cmax of DLM was comparable between children and adults. After a single oral dose of DLM (0.1 mg/kg), the predicted Cmax was 1.4, 1.7, 2.4, and 2.0 pg/g, respectively, for children ages 1, 5, and 10 years old, and adults. The observed species differences in age-dependent PK seems to be largely attributable to the species differences in enzyme ontogeny and resulting metabolic clearance rates for DLM as well as differences in doses. In conjunction with age-specific exposure data, the current model will enable us to evaluate early life sensitivity in humans under environmentally relevant exposure conditions for this chemical and can be readily applicable to other pyrethroids with proper parameterization (supported by CAPHRA).


Ethanol (EtOH), which induces fetal alcohol spectrum disorder-related effects, was selected to investigate the use of PBPK models in a quantitative in vivo to in vitro extrapolation (IVIVE) paradigm. Mouse whole embryo culture data on EtOH time-concentration dependent induction of neural tube defects (NTDs) was used to assess embryotoxicity. Initially, published rat life-stage PBPK models were extrapolated to adult non-pregnant and pregnant mice and humans using species specific (SS) tile equations and parameters. Each model was able to predict blood EtOH concentrations (BECD) following multiple routes of exposure. Once calibrated with published data, the mouse and human models produced adequate simulations (within factor of 2) of time-course BEC data. An IVIVE was performed by estimating SS BECDs in rat and mouse dams using in silico simulations for induction of NTDs in vitro. Then, a human model IVIVE was performed using rodent in vitro and pregnant human toxicity and epidemiological reports of EtOH exposure during windows of gestational susceptibility. As reported in the human literature, 150 mg/dl BECD is associated with developmental neurotoxicity. In human embryonic stem cells (hESCs), 145 mg/dl EtOH is associated with apoptosis and altered neuroprogenitor differentiation and proliferation. The similarity in effective concentration illustrates the utility of using BECD in pregnant women as a predictor of adverse developmental effects. However, since hESCs are more sensitive to EtOH-induced effects than mouse conceptsus in vitro, a human IVIVE based on mouse in vitro data exceeds the reported 150 mg/dl BECD. The human model predicted exposure concentrations within a factor of 2 of the observed toxic BECDs. This research demonstrates both the capability of models to extrapolate between in vitro animal and in vivo human studies and the first pregnancy models for EtOH in mice and humans. The views expressed in this abstract are those of the authors and do not necessarily represent the views or policies of the U.S. EPA.


Neurodegeneration is the underlying cause of a vast majority of neurological disorders and often a result of brain trauma, stroke, or neurotoxic insult. A widely used and reliable method for labeling degenerative neurons in ex vivo brain tissue involves the use of the Fluoro-Jade stains. Recently we described a novel method for...
the labeling of degenerating neurons in unfixed brain tissue samples using Fluoro-Jade-C (FJC) (Gu et al. Journal of Neuroscience Methods 2012, 208:40-43). This method is simple, fast, and applicable to unfixed brain tissue sections and works at neutral pH. Based on these features, we extended our experimentation by examining the utility of FJC in vitro using cell cultures. Using neural stem cells, for example, we report here specific FJC labeling following treatment with neurotoxins such as cadmium. The FJC labeling appears to be specific to neural cells, since other cell types such as cultured kidney epithelial cells did not show similar labeling even when cells were dying or dead. Further characterization and validation of this in vitro approach is underway. By employing FJC labeling in multi-well culture plates and using high-content time-lapsed recordings and additional techniques, a primary goal will be to achieve high-throughput monitoring and analysis of morphological, biochemical and molecular changes associated with the entire neurodegenerative process in cell culture models after neurotoxicant exposures. This in vitro approach has the potential to not only reduce animal use and suffering in toxicity tests but also to facilitate high-throughput screens for potentially neurotoxic compounds.

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900 Evaluation of the Neuroactivity of ToxCast Compounds Using Multiewell Microelectrode Array Recordings of Primary Cortical Neurons.

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Assessment of spontaneous activity in neuronal cultures on microelectrode arrays (MEA) is a sensitive method to detect responses to drugs, chemicals, and particles. While single-well MEA systems lack the efficiency to screen large numbers of compounds, recently developed multi-well MEA systems have increased throughput of MEAs. The present experiments examined the ability of a subset of EPA’s ToxCast library of compounds to alter neuronal activity using 48 well MEA plates. Sixty-eight compounds were selected from the ToxCast Phase I and II libraries based on known neuroactivity or data from 14 ToxCast in vitro assays indicating that they interacted with ion channels. Two compounds expected not to alter neuronal function, acetylcholine and glutamate, were included as negative controls. One hr of baseline activity was recorded prior to exposing the cortical networks to 40 μM of each compound for 1 hr and the weighted mean firing rate (wMFR) was determined in the absence and presence of each chemical. All experiments were conducted on day in vitro 14 or 15. Based on DMSO-treated wells, chemicals that increased or decreased activity by >25% were considered hits. Of the 68 compounds, 47 altered wMFR by more than the threshold. Saccharin and acetylamo- phile change wMFR beyond the threshold. Four of six pyrethroids, and 3/3 conazoles were detected as hits. Interestingly only 1/7 nicotinic agonists (nicotinic acetylcholine receptors) were hits, but were close to the threshold. These data demonstrate that multiwell MEAs can be used efficiently to screen chemicals for potential neurotoxicity and that the results are concordant with predictions from the ToxCast in vitro assays. To further study the effects of TBBPA on calcium-permeable nACh receptors, effects in B35 cell line were examined using single cell fluorescent calcium imaging. TBBPA (≥1 μM) inhibits ACh receptors in B35 cells as evidenced by a reduction in the ACh-evoked increases in the intracellular calcium concentration (Ca2+). Additionally, TBBPA (≥1 μM) induces a strong and concentration-dependent increase in basal [Ca2+]i in B35 cells. In dopamineergic PC12 cells, TBBPA (≥1 μM) also increases basal [Ca2+]i. The increase in basal [Ca2+]i is also evident under calcium-free conditions, indicating it originates from intracellular calcium stores. Moreover, depolarization-evoked increases in [Ca2+]i are strongly reduced by TBBPA (≥1 μM), indicating TBBPA-inhibited induction of voltage-gated calcium channels. Our in vitro studies thus demonstrate that TBBPA exerts multiple adverse effects on functional endpoints for neurotransmission, justifying the quest for flame retardants with reduced neurotoxic potential. Funding: EU FP7 (ENVIRO; grant agreement FP7-ENV-2008-1-226563).

Tetramethylbisphephol-A (TBBPA) is a widely used brominated flame retardant. Studies in the in vitro neurotoxic potential of TBBPA focused on cytotoxicity and presynaptic effects of neurotransmission, while recent studies indicate that persistent organic pollutants can also affect postsynaptic inhibitory human GABAA and excitatory glycine receptors. Possible effects of TBBPA on these neurotransmitter receptors are of considerable interest as these receptors are critically involved in neurotransmission, synaptic plasticity and brain development. We therefore investigated the effects of TBBPA on these receptors, expressed in Xenopus oocytes, using the two-electrode voltage-clamp technique. Our results demonstrate that TBBPA acts as full (≥10 μM) and partial (≥0.1 μM) agonist on human GABAA receptors, while it acts as antagonist (≥10 μM) on human γ4 nACh receptors. To further study the effects of TBBPA on calcium-permeable nACh receptors, effects in B35 cell line were examined using single cell fluorescent calcium imaging. TBBPA (≥1 μM) inhibits ACh receptors in B35 cells as evidenced by a reduction in the ACh-evoked increases in the intracellular calcium concentration ([Ca2+]i). Additionally, TBBPA (≥1 μM) induces a strong and concentration-dependent increase in basal [Ca2+]i in B35 cells. In dopaminergic PC12 cells, TBBPA (≥1 μM) also increases basal [Ca2+]i. The increase in basal [Ca2+]i is also evident under calcium-free conditions, indicating it originates from intracellular calcium stores. Moreover, depolarization-evoked increases in [Ca2+]i are strongly reduced by TBBPA (≥1 μM), indicating TBBPA-inhibited induction of voltage-gated calcium channels. Our in vitro studies thus demonstrate that TBBPA exerts multiple adverse effects on functional endpoints for neurotransmission, justifying the quest for flame retardants with reduced neurotoxic potential. Funding: EU FP7 (ENVIRO; grant agreement FP7-ENV-2008-1-226563).

902 In Vitro Neurotoxicity of Tetramethylbisphephol-A (TBBPA).


Seizurogenic neurotoxicity produces significant drug attrition during drug discovery. Current available in vitro assays fail to predict this toxicity due to the failure of general cytotoxicity assays to predict sublethal target specific electrophysiological liabilities. Ion channel and receptor activity assays can be used to predict some seizure potential, but this only focuses on specifically measured targets for prediction and may miss a response which relies on a combination of endpoints. Most evaluation of seizure inducing compounds occurs later in preclinical development in in vivo studies which have higher costs and could result in species specific results. Therefore, the development of an in vitro assay to screen compounds for seizurogenic potential in a human neural model would give the potential to screen compounds earlier at lower cost and greater reliability. Here we demonstrate the use of multiwell MEAs to screen for seizurogenic compounds in human IPS derived neurons (hSNs). hSNs were seeded in 48 well MEAs and spontaneous activity began at 3 days post plating with greater activity at 7 days when the assay is performed. Neural action potentials were detected and the results were reported as mean firing rate (MFR). The seizurogenic compounds tested show dose dependent increase in MFR with changes in spike train organization, while all of the negative controls were unaffected. The seizurogenic compounds tested were PicROTOX, gabazine, L-Glutamate, and pentyleneetetrazol (PTZ). Negative compounds tested were acetaminophen, naproxen, and DMSO. To further demonstrate the responsiveness of the cells in the assay, we tested Domoic Acid, a neurotransmitter known to cause amnesia, and found that it completely blocked network activity while not causing cell death. These results illustrate the power of the human Neural MEA assay for predicting compound induced neural toxicity, especially the seizurogenic response.

903 Modulation of Nicotinic Acetylcholine Receptor by Brominated and Alternative Halogen-Free Flame Retardants.


The large scale use of brominated flame retardants (BFRs) is associated with ecological and toxicological concerns. Previous in vitro research demonstrated that nicotinic acetylcholine (nACh) receptors are a direct target for e.g., PBDEs and TBBPA. These (neuro)toxic effects of BFRs argue for replacement by safer and less persistent alternatives. Since it is essential to assess the (neuro)toxic potential of proposed halogen-free flame retardants (HFFRs) before they are used on large scale and in high volume, we measured the effects of three frequently used BFRs and 13 possible halogen-free substitutes on the function of human γ4β2 nACh receptors, expressed in Xenopus oocytes, using the two-electrode voltage-clamp technique. Our initial rank-order potency based on the in vitro inhibition of nACh receptors indicates the neurotoxic potential of the HFFR triphenylphosphinite (TPP), aluminium diethylphosphinate (Alpi), ammonium polyphosphate (APP), and the nanoclay cloisite 30B (MMT). However, additional studies focusing on expected concentrations in humans and the environment are required before these compounds can be excluded as viable alternatives. Importantly, five out of the sixteen tested compounds (brominated polystyrene (BPS), bispheno-A bis(diphenylphosphate) (BDP), resorcinoil bis(diphenylphosphate) (RDP), 9,10-dihydro-9-oxa-10- phosphaphenanthrene-10-oxide (DOPO), and zinc stannate (ZS) are classified as not potent. Based on this specific neurotoxic endpoint, these five compounds could therefore be selected for additional testing to further assess the viability of these HFFRs as alternatives to replace current BFRs.
904 Neurotoxic Effects of Bisphenol AF by the Calcium-Induced ROS and MAPKs.
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Bisphenol AF (BPAF), a newly introduced chemical structurally related to bisphenol A (BPA), is used extensively in fluoroelastomers and polyesters, and has been known to induce estrogen-dependent responses. However, the toxicity of BPAF is largely unknown except for its endocrine-related effects. In this study, we investigated the neurotoxicity of BPAF and underlying mechanisms of action using hippocampal cell line (HT-22) and mouse primary neuronal cells. We found that BPAF induced apoptosis in both HT-22 and primary neuronal cells. To clarify the underlying mechanisms of BPAF-induced apoptosis, various signaling molecules were evaluated. BPAF increased the level of intracellular calcium, followed by the generation of reactive oxygen species (ROS). BPAF upregulated the phosphorylation of mitogen-activated protein kinase (MAPK): extracellular signal-regulated kinase (ERK), p38 and c-Jun-N-terminal kinase (JNK), and nuclear translocation of nuclear factor (NF)-κB. Using specific inhibitors, we confirmed that calcium, ROS, p38, and JNK mediated the BPAF-induced apoptosis. In addition, BPAF inhibited microglial activation in a microglia/neuroblastoma coculture model by the reduction of nitric oxide production. We found that BPAF interrupted the normal physiological functions of microglia at non-toxic levels. Taken together, our results suggest that BPAF, the substitute of BPA, also have neurotoxic properties.

905 Potent Induction of a Series of Endogenous Antioxidative Enzymes by Triterpenoid CDDO-IM Leads to Neuprotection Against Oxidative and Electrophilic Injury.
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Evidence suggests oxidative and electrophilic stress as a major factor contributing to the neuronal cell death in neurodegenerative disorders, especially Parkinson’s disease (PD). Early depletion in the levels of thiol antioxidant glutathione (GSH), which may lead to generation of reactive oxygen species, is an important biochemical feature of PD. However, whether induction of endogenous antioxidative enzymes by a novel triterpenoid CDDO-IM (2-cyano-3,12-dioxooleana-1,9(11)-dien-28-oic acid) affords protection against oxidative and electrophilic neurotoxicity has not been carefully investigated. Retinoic acid-induced differentiation of human neuroblastoma SH-SY5Y cells are known to possess properties of mature neurons and thus have been widely used in vitro model for the study of neurotoxicity and neuroprotection. In this study, we showed that incubation of retinoic acid-induced differentiation of human neuroblastoma SH-SY5Y cells with nanomolar concentrations of CDDO-IM (1-400 nM) for 24 h resulted in significant increase in the levels of reduced glutathione (GSH) and NAD(P)H:quinone oxidoreductase 1 (NQO1), two critical cellular defenses in detoxification of reactive oxygen species and electrophilic quinone molecules. Pretreatment of the cells with CDDO-IM was found to afford remarkable protection against the neurotoxicity elicited by acrolein, 4-hydroxyxynonenal, 3-morpholinosynonimine hydrochloride, xanthine oxidase/xanthine, and hydrogen peroxide. Taken together, this study demonstrates for the first time that CDDO-IM potently induces the cellular GSH system and NQO1 in retinoic acid-induced differentiation of human neuroblastoma SH-SY5Y cells, which is accompanied by dramatically increased resistance of these cells to the damage induced by various neurotoxins. The results of this study may have important implications for the development of novel neuroprotective strategies.

906 1,3-Dinitrobenzene Induces Age-Specific Responses in Primary Neurons.
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During the normal aging process, the brain successively loses the ability to cope with pathophysiologic stressors. The accumulation of iron in the aging brain potentially contributes to this by increasing redox activity. Astrocytes participate in the preservation of neurons during chemically induced stress by releasing adenosine, thereby silencing the electrical (hence energetic and metabolic) requirements of neurons. 1,3-dinitrobenzene (DNB) stimulates adenosine release from astrocytes, and is used as a neural stress probe in this study. Primrose stress and astrocytes were exposed to DNB (0, 100nM, or 10 μM) and 10 μM adenosine with 100nM or 100μM DNB. Neurons were isolated from male F344 rats (1 mo, 3 mo, & 18 mo), maintained for two weeks and exposed to DNB. Astrocytes were isolated and maintained for five weeks and cocultured with neurons. Adenosine triphosphate (ATP) was measured in both cell types. Neurons from 3 mo brains exhibited higher ATP than neurons from other age groups. Cocultured astrocytes had higher ATP levels than neurons at all age groups. Neurons were assayed at a later timepoint for lactate dehydrogenase (LDH). Neurons from brains of 1 mo and 18 mo rats, but not 3 mo rats, exposed to DNB exhibit statistically significantly higher LDH than neurons treated with both DNB and adenosine. Immunocytochemistry for the presence of transferrin (TF) and mitochondrial frataxin (mtFr) was performed on neurons. Increases in TF were observed in older neurons. Colocalization of TF and mtFr was observed in the oldest age group. These results suggest that neurons from older animals sequester more mitochondrial iron than younger cohorts, increasing potential for higher redox activity, rendering them more vulnerable to toxic insults such as DNB. Future work will elucidate the significance of mitochondrial iron in the aging process.

907 Magnetic Resonance Imaging and Spectroscopic Markers of Kainic Acid-Induced Excitotoxicity in the Rat Brain.
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Neurotoxicity assessment in drug development is typically accomplished using microscopic analysis, which may be time consuming and not always comprehensive. In this study we describe changes in brain after acute administration of kainic acid (KA) using non-invasive magnetic resonance imaging (MRI) and spectroscopy (MRS) in comparison to histopathology. Adult male Sprague-Dawley rats (N = 24, 361 ± 36 g) were anesthetized with isoflurane and positioned inside a 7 tesla MRI scanner. T2 relaxation mapping of the whole brain and proton MRS in the left hippocampus were performed. KA (10 mg/kg, ip) was then administered to all animals after baseline scans and imaging was continued for another 2 hours. One group of animals (N = 6) was euthanized at this time. The MRI procedure was repeated one day after treatment (N = 6) and 2 days later (N = 12) and these animals were euthanized. All rats were perfusion-fixed and their brains underwent histopathological assessment. KA led to an increase in T2 in the hippocampus as early as 1 hour after administration. These changes were more pronounced at 2 hours and drastically so at 1 and 2 days after the treatment at which point the findings spread to wider areas of the brain, including the amygdala, thalamus, and cortex. MRS revealed immediate increases in glutamate and glutamine concentrations right after KA administration followed by decreases at 1 and 2 days, at which point N-acetyl-aspartate signal was also decreased. Lactate was not detectable in the normal brain but appeared at 15 min and increased to a maximum at 2 hours after treatment. At 1 and 2 days lactate was still detectable. The T2 map correlated with histopathological changes in the brain at all time points. These data provide the basis for the development of imaging biomarkers for the early detection of neurotoxicity, which would benefit public health by increasing the number of tools available for safety evaluation of new drugs. (Supported by NCTR and CDER, FDA, #E0714801).

908 Effect of Acute Administration of Beta-N-Methylaminoalanine (L-BMAA) on Rat Hippocampus Neurochemical Profile Determined with 1H Magnetic Resonance Spectroscopy (MRS).
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L-BMAA is a non-proteogenic amino acid present in cyanobacteria and in cycad seed, exposure to which is implicated in western Pacific amyotrophic lateral sclerosis and parkinsonism-dementia complex (ALS-PDC). After absorption, L-BMAA is transported into the brain, metabolized, and L-BMAA and/or its metabolites may accumulate in brain protein. This bound form may serve as a sink from which L-BMAA or its metabolites are released, contributing to its neurotoxic effect. L-BMAA has been shown to be excitotoxic, acting as a glutamate receptor agonist. In
this study we used non-invasive 1H-MRS to investigate the changes in the hip-
cocampal neurometabolitic profile after single dose of L-BMAA. Nine adult male
Sprague-Dawley rats (390 ± 8 g) were anesthetized with isoflurane and positioned
in 7 tesla MRI scanner. MRS in the left hippocampus (voxel size 4 x 4 x 2 mm) was
performed before and for 1.5 hrs after BMAA administration (100 mg/kg, ip).
Signal intensity of N-acetyl-aspartate, gamma-aminobuteric acid (GABA), creatine,
choline, glutamate, glutamine, myo-inositol, and taurine were measured relative to
water peak in water-unsuppressed reference spectrum. At the end of observation pe-
toid the GABA signal was elevated relative to the baseline (+23%, P = 0.051) and
taurine was decreased (-10%, P = 0.014). Signal intensity of other neurotransmit-
tors did not change. These data may be consistent with L-BMAA mediated gluta-
mate agonist activity. Both GABA and taurine are released into extracellular space
after stimulation of NMDA receptors. Increase in GABA may be due to a combi-
nation of uptake and increased synthesis. The lowering of hippocampal taurine may
reflect a reduction in re-uptake after release. These data suggest that L-BMAA has
measurable effect on glutameric pathways in undisturbed, intact tissues in rats.

909 Pattern of Pathologic Changes Observed in the CNS of
Chronic Toluene Abusers.

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Toluene is a common volatile organic chemical (VOC) found in petroleum-based
products and has wide-spread industrial use as a solvent in paints, lacquers, inks,
adhesives and cleaning agents (decreasing). The toxicity of toluene has been well
described in experimental animals, occupationally exposed humans and healthy
volunteers. Considerable toxicological data has also been obtained from individuals
who purposely expose themselves to high levels of toluene and/or toluene contain-
ing solvents to achieve the narcotic effect common to most organic solvents.
Collectively, this data has demonstrated that high-dose prolonged exposure to
toluene can cause neurological toxicity, ranging from non-specific, reversible symp-
toms to permanent pathological changes in the white matter of the brain. Evalua-
tion of chronic toluene abusers by magnetic resonance imaging (MRI) have
demonstrated a series of pathologic alterations that appear to be common to most, if not
all, toluene abusers that suffer from permanent neurological alterations. These in-
dividuals almost always provide evidence of direct toxic insult, including diffuse atro-
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911 Nitrosative Stress in Chemococonvulant-Induced
Epileptogenesis.

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Reactive oxygen and nitrogen species (ROS/RNS) are mediators of oxidative stress
and are increased in the brain as a result of seizure activity and reactive gliosis; yet
their contributing role in the process by which injury leads to epilepsy (i.e. epilep-
togenesis) is largely unknown. Our previous work in the kainate model of chemo-
convulant-induced epilepsy has confirmed significant oxidative macromolecular
damage to hippocampal mitochondria (Ryan et al 2012). The primary goals of this
study were to assess the consequences of nitrosative stress in cellular compartments
to gain greater mechanistic insight into mitochondrial dysfunction during epilepto-
genesis. Rats were exposed to a single high dose of kainate (11mg/kg) or vehicle
and monitored by video and/or EEG for seizure activity 6 hrs. Nitrosative stress
was measured in the susceptible hippocampal brain region acutely after kainate ad-
ministration (8hr, 24hr, and 48hr), 1wk after kainate prior to development of
epilepsy (latent period) and during the chronic stages of epilepsy (3wk and 6wk).
Analysis from frozen hippocampal sections indicated global astrogliosis (GFAP)
and microgliosis (Iba-1) after kainate. Hippocampal nitric oxide spiked 2-fold 8hr
after kainate and remained elevated throughout epileptogenesis. Preliminary results
suggest this could be associated with increased iNOS detected by immunohisto-
chemical staining. Furthermore, cellular and mitochondrial 3-nitrotyrosine (3NT)
progressively and significantly increased during epileptogenesis indicating protein
nitration and subsequent immunofluorescent co-labeling showed that 3NT is local-
ized to hippocampal neurons (NeuN). This study demonstrates a probable mecha-
nism for gliosis-mediated nitric oxide production and downstream protein damage
to hippocampal neurons in epileptogenesis. These studies propose a role for target-
ing oxidative and nitrosative species to delay or prevent disease progression by
inhibiting detrimental protein nutrition. NS0395978 (MP) and NS R21 NS0572099
(M.P.)

912 Ozone-Induced Changes in Oxidative Stress Parameters
in Brains of Adult, Middle-Age, and Senescent Brown
Norway Rats.

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Understanding life-stage susceptibility is a critical part of community based human
health risk assessment following chemical exposure. Recently there is growing con-
cern over a common air pollutant, ozone (O3), and adverse health effects including
dysfunction of the pulmonary, cardiac, and nervous systems. Oxidative stress (OS)
is a known contributor in multiple organ toxicities and plays an important role in
age-related diseases. Growing evidence implicates OS in O3 toxicity. The current
study explored OS as a potential toxicity pathway for O3 exposure and addressed
whether these effects are life stage-dependent. OS-related measures included reac-
tive oxygen species (ROS) production [NADPH Quinone oxidoreductace 1 (NQO1),
NADH Ubiquinone reductase (UBIQ), antioxidant homeostasis (total antioxidiant
substances (TA), superoxide dimuastase (SOD), γ-glutamylcysteine synth-
ethe (γ-GCS)], and oxidative damage (total aconitase). Male Brown Norway rats
(8, 14, and 24 months) were exposed to O3 (0.25 or 1 ppm) via inhalation for 6
h/day, 2 days per week for 13 weeks. Frontal cortex (FC), cerebellum (CB), stria-
tum (ST), and hippocampus (HP) were dissected 24 hours after last exposure,
quick frozen, and stored at 80°C until analysis. Results indicated life stage-related
increases in ROS production (~ 2x in UBIQ and ~ 1.5x in NQO1 in striatum), slight
decreases in antioxidant homeostatic mechanisms (TAS, γ-GCS, and SOD), and a
decrease in aconitase activity. The effects of O3 exposure were brain area-spe-
cific, with the striatum being more sensitive than other brain regions. With regard
to life stage, the effects of O3 appeared to be greater in 4 month old than 12 or 24
month old rats. These results suggest that the Oxidative Stress Pathway for O3,
but the complex interactions between age, exposure and brain region require
further investigation. (This abstract does not necessarily reflect USEPA policy).
The Neurotoxic Effects of Tributyltin on Tokai High Avoider (THA) Rats Evaluated by the Sidman Electric Shock Avoidance Test.


Tributyltin (TBT) compounds have been known as environmental pollutants. Although neurotoxicity is one of the major toxicities of TBT, the effects of TBT on learning ability have not yet been clearly demonstrated. Wistar-derived Tokai High Avoider (THA) rats, which were developed at Tokai University, can achieve stable high-level avoidance ability and small individual differences. We evaluated the neurotoxicity of TBT in THA rats by Sidman electric shock avoidance test. The male and female THA rats were administered TBT in their food at 0 and 125 ppm after weaning (males, n = 5/group; females n = 4/group). From 6 weeks of age, the Sidman test sessions were performed for 60 min/day for 10 consecutive days. Rats can avoid electric shocks by pressing a lever. Avoidance rates for shock exposures were calculated for the first and second halves (30 minutes each) of each session. The mean values of the avoidance rates were compared between the control and TBT groups for each sex, and those of the TBT-exposed group were lower than those of the control for both sexes for all the sessions. Significantly lower mean values of the avoidance rates compared to the control were observed in the first 30 minutes of the sessions in male rats on days 2 and 5. It is suggested that learning ability was impaired in the THA rats exposed to TBT at 125 ppm. To detect the statistically significant differences in future studies, more THA rats should be used.

Protein Biomarker Panel of Cisplatin-Induced Neurotoxicity in a Preclinical Model.


Background: Neurotoxic brain damage is a widely recognized adverse effect of chemotherapy affecting overall outcome and quality of life cancer survivors. Effective clinical monitoring of neurotoxicity using simple and reliable assays could provide timely information to clinicians allowing them to adjust treatment and reduce neurologic and cognitive side effects of chemotherapy. The goal of this study was to investigate spatiotemporal distributions in the brain, CSF and serum of glial, neuronal and inflammatory biomarkers: ubiquitin C-terminal hydrolyase-1 (UCH-L1), glial fibrillary acidic protein (GFAP), cell-spectrin break down products (SBDPs), microtubule-associated protein 2 (MAP2), myelin basic protein (MBP), MBP break down products (MBP-BDP), intercellular adhesion molecule (ICAM) and their relationships to cisplatin-induced neuropathology in rats.

Methods: Neurotoxicity in adult rats was induced by cisplatin (10 mg/kg, i.p.). The levels and localization of biomarkers in the brain were examined by immunohistochemistry (IHC) and their levels in CSF and serum were evaluated by ELISA. Results: IHC revealed that cisplatin caused abnormal changes in the brain starting at 6 h and increasing at 24 and 48 h including gliosis, determined by increased GFAP level, dendritic damage determined by decreased MAP2 level, neuronal demyelination determined by increased level of MBP-BDP, and inflammation determined by increased level of ICAM. Serum levels of UCH-L1, GFAP and SBDP150 were significantly increased at 24 h after cisplatin administration as compared to controls. The levels of these biomarkers were correlated with IHC brain pathologies and survival.

Conclusion: This study demonstrated the potential of using levels of glial and neuronal proteins in blood for assessment of cisplatin induced neurotoxicity. In addition, assessment of serum levels of biomarkers can provide information on underlying mechanisms of neurotoxicity and facilitate a personalized treatment to minimize side effects during and after chemotherapy.
Dopaminergic nuclei within the basal ganglia are important for control of motor function but are highly sensitive to damage from oxidative stress, inflammation, and environmental neurotoxins. Here we propose that inhibition of transmitter-evoked calcium (Ca2+) signaling in astrocytes may contribute to this sensitivity because ATP-dependent Ca2+ waves in these cells modulate diverse trophic functions in the CNS, including metabolism, synaptic activity, and regional cerebral blood flow. To examine mechanisms underlying alterations in Ca2+ signaling in astrocytes, we postulated that dopaminergic neurotoxins of the basal midbrain would acutely inhibit ATP-induced Ca2+ signals in astrocytes through a mitochondrial cascade. To test this hypothesis, we examined the capacity of MPP+ and 6-Hydroxydopamine (6-OHDA) to block ATP-dependent Ca2+ waves and transients in primary striatal astrocytes. Calcium imaging studies revealed a dose-dependent decrease in ATP-induced intracellular Ca2+ transients and mechanically stimulated Ca2+ waves following acute application of both MPP+ and 6-OHDA. These compounds inhibited mitochondrial Ca2+ influx, suggesting the transient receptor potential channel, TRPC3, as a probable target. MPP+ inhibited OAG-induced intracellular Ca2+ transients similarly to the TRPC3 channel antagonist, pyrazole-3, whereas 6-OHDA only partly suppressed OAG-induced Ca2+ transients. In whole cell patch clamp experiments conducted in TRPC3-overexpressing cells, acute application of MPP+ completely blocked TRPC3-like currents, whereas only partial inhibition was detected in the presence of 6-OHDA. These findings indicate that dopaminergic neurotoxins that have cationic properties differentially inhibit TRPC3 in astrocytes, thereby diminishing the amplitude of ATP-induced Ca2+ transients. These compounds may therefore share a common mechanism of toxicity in their capacity to acutely disrupt trophic trophic functions reliant on this signaling mechanism.
measurable neurobehavioral endpoints was evaluated. Socially defeated or submis- sive rats exhibit impaired exploratory behavior relative to social- ly dominant rats. It was hypothesized that any impact of social dominance on a neurobehavioral endpoint (e.g., motor activity) would correlate with a difference in bodyweight gain between the two pair-housed rats. However, the control data from five 90-day studies (N=114) indicated no significant correlation between the differ- ence in bodyweight gain within each cage and the difference in motor activity, grip strength, hindlimb splay, or body temperature values within the same cages. Our second objective was to optimize the use of a photo-beam-based system which computes multiple parameters for motor activity assessment and is widely used in the industry. We evaluated baseline data from eight studies (N=816), to select two re- presentative motor activity parameters from a list of multiple options. Duration of movement and number of ambulatory movements offered the greatest unique value (lowest correlation coefficients) and were the best predictors (highest R-squared val- ues) of the remaining motor activity parameters. A positive control study con- ducted with carbaryl and amphetamine using pair housing conditions demon- strated neurobehavioral effects that were consistent with previously published literature. Based on the analysis of data from multiple studies including a positive control, we concluded that pair housing does not interfere with neurobehavioral evaluations and that motor activity can be characterized effectively using duration of movement and number of ambulatory movements in a photo-beam-based sys- tem.

In conclusion, this validation study demonstrated the feasibility of EEG recordings in chair-restrained cynomolgus monkeys since sedating as well as seizure-inducing effects could be demonstrated.
MicroRNAs are promising noninvasive biomarkers of drug-induced toxicities due to their tissue-selective expression, rapid release post-injury, and stability in biofluids, but their low levels in biofluids present challenges to their reliable quantification. The HESI Genomics committee initiated a collaborative study to assess best practices for measuring injury-related microRNAs in biofluids. Samples were derived from a model of acute cardiotoxicity in male Wistar rats induced by a single s.c. injection of 0.5 mg/kg isoproterenol. The heart-enriched microRNAs miR-208, miR-499, and miR-1 increased by approximately 10-fold in serum and plasma 4 hr after treatment. In a follow-up study using the same model system, urine was collected overnight and plasma at 24 hr post-injection. Biofluids were pooled from 5 rats per group and aliquots were sent to 10 laboratories for analysis of levels of the 3 heart-enriched microRNAs. At each site, samples were assayed using a standard protocol and the data normalized to levels of a spiked-in ath-miR159a control. The results from this interlaboratory analysis of multiple preanalytical and technical issues provide guidelines for the accurate measurement and reporting of injury-related microRNAs in biofluids.

MicroRNAs (miRNAs) are small, conserved, non-coding RNAs that modulate gene expression post-transcriptionally. They are remarkably stable in various body fluids like urine and thus suggested as new non-invasive biomarkers (BM) of tissue injury, especially since some miRNAs are produced in cell- or tissue-specific manners offering the possibility to identify the site of injury. Furthermore, their tissue expression profiles may be used for investigation of the mechanistic details of compound-induced toxicities. We investigated miRNAs in urine of male Crl:WI (Han) rats rats treated with nephrotoxins: Cisplatin (Cp), to specifically elicit proximal tubular injury; and male Wistar Kyoto rats treated with a nephrotoxic serum (NTS) containing antibodies against the glomerular basement membrane to induce glomerular damage. The expected tissue injury was confirmed by histopathology. Total RNA including small RNAs was isolated from urine on day 3, 5, 8, 15 and 26 after a single dose of Cp (0, 1 and 3 mg/kg) and on day 8 and 14 after NTS injection (0.1, 0.25 and 0.5 mL/100g bw). Urinary miRNA profiles were measured with TaqMan® MicroRNA Card (Tldas). miRNA levels were normalized to a non-endogenous spiked-in control miRNA and to urinary creatinine. Comparing Cp-high dose vs. control samples and NTS-high dose vs. control samples we identified 131 and 68 miRNA, respectively, with significantly increased urinary levels. About 65 miRNAs were affected by both treatments, whereas app.10 and 35 urinary miRNAs were exclusively affected by NTS and Cp treatment, respectively. Furthermore, TDLA analysis of kidneys from NTS-treated rats showed deregulation of many miRNAs upon glomerular injury. Our results suggest that urinary miRNAs may be used as promising new and site-specific BMs for nephrotoxicity and encourage further investigations of their role in different types of kidney injury.
For the TLDA–derived PCR data, a modified ΔΔCt method was applied: delta-Ct values obtained by subtracting ath-miR-159a Cts from target miRNA Cts were de-
logarithimized and divided by the corresponding uCrea value. This normalization reduced the variation of the delta-Ct values for miRNAs across samples. Then ra-
tios were calculated between treated and time-matched control groups. With this method we identified 131 miRNAs with significantly increased urinary levels on days 3 and 5. Twenty miRNAs with distinct time-dependent profiles were then measured using TaqMan® MicroRNA Assays. The data were analysed either with the modified ΔΔCt method, or absolutely quantified using synthetic miRNA stan-
dard curves run in parallel. With both methods we obtained comparable results with respect to observed direction of changes and time course of affected urinary miRNAs.

Our results indicate that urinary miRNAs may be used as BMs for nephrotoxicity in rats. Furthermore, normalization to uCrea, which is recommended for urinary BMs, ap-
pears feasible for analysis based either on the PCR-generated Ct values only, or in-
cluding absolute quantification.

Consistent differentiation of embryonic, or induced pluripotent, stem cells into functional hepatocyte-like cells available on demand for hepatotoxicity screening would be a valuable resource for drug development programs. Assessments of stem cell-derived hepatocyte-like cell maturity often lack quantitative comparisons with primary human hepatocytes and the presumed fetal-like phenotype currently achieved is concluded from observations of cell morphology and the detection of a small number of proteins such as albumin, alpha fetoprotein and alpha 1-antit-
rypsin. The identification of a hepatocyte-specific marker that can be quantified ab-
solutely would aid the assessment of cells and be useful for inter-laboratory com-
parisons. MiR-122 is a micro-RNA species that is very highly enriched in hepaticytes. We hypothesised that this molecule would be a useful biomarker of differen-
tiation and could also serve as a more sensitive indicator of hepatocyte-spe-
cific injury. Our study demonstrates that miR-122 is a specific and sensitive marker of hepatocyte phenotype, present at high levels in primary human hepatocytes and in fetal liver samples but of low abundance or absent in hepatoma and non-hepatic human cell lines. Expression of miR-122 was greatly elevated in stem cell-derived hepatocyte-like cells compared to the undifferentiated stem cell lines. Furthermore we demonstrate miR-122’s utility as a sensitive marker of hepatotoxicity in primary hepatoctyes and a potential adjunct to screens such as lactate dehydrogenase release and ATP depletion. We conclude that quantitative measurement of miR-122 may be used as sensitive and specific marker during the hepatoctyte differentiation process or to confirm the hepatocyte-like status of stem cell-derived material fol-
lowing hepatotoxicity screening assays.

Exposure to lead (Pb) causes numerous deleterious health effects including devel-
opmental and cognitive abnormalities in children. Early detection of lead exposure is critical to reduce serious health problems arising from chronic lead toxicity. Circulating nucleic acid based biomarkers are present in various bodily fluids such as blood, serum, and urine. A number of studies have reported the use of circulat-
ing nucleic acids as biomarkers for pathological conditions and developmental stages. In this study, a Long Evans rat model system is used to identify lead expo-
sure-induced circulating nucleic acid biomarkers in peripheral blood. Two different groups of animals are used, females and time-pregnant Long Evans rats.

Furthermore normalization to uCrea, which is recommended for urinary BMs,
appears feasible for analysis based either on the PCR-generated Ct values only, or in-
cluding absolute quantification. Our results indicate that urinary miRNAs may be used as BMs for nephrotoxicity in rats. Furthermore, normalization to uCrea, which is recommended for urinary BMs, appears feasible for analysis based either on the PCR-generated Ct values only, or including absolute quantification.

Consistent differentiation of embryonic, or induced pluripotent, stem cells into functional hepatocyte-like cells available on demand for hepatotoxicity screening would be a valuable resource for drug development programs. Assessments of stem cell-derived hepatocyte-like cell maturity often lack quantitative comparisons with primary human hepatocytes and the presumed fetal-like phenotype currently achieved is concluded from observations of cell morphology and the detection of a small number of proteins such as albumin, alpha fetoprotein and alpha 1-antit-
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lowing hepatotoxicity screening assays.

The present study investigated the role of oxidative stress as a possible risk factor for colorectal cancer in humans. Blood samples were collected from 232 subjects in-
cluding 102 healthy controls, 90 patients with polyps, 39 patients diagnosed with colorectal cancer. We measured both direct and oxidative DNA damage in periph-
eral white blood cells (WBRCs) using alkaline (direct DNA damage) and formami-
dopyrimidine DNA glycosylase modified (oxidative DNA damage) Comet assay. Trolax equivalent total antioxidant capacity (TEAC) in plasma was also determined spectrophotometrically. Results showed that Oxidative Stress induced DNA dam-
age was associated with a significantly higher probability of colorectal cancer in males (p < 0.008) but not in females (p = 0.675) compared to healthy controls. Direct DNA damage also showed a marginally significant effect on colorectal can-
cer (p = 0.071). Results from the TEAC assay showed a positive trend on colorectal cancer, but was not statistically significant (p = 0.230). Oxidative stress induced DNA damage, TEAC, and direct DNA damage did not show significant relationships with probability of polyps. Females showed significantly higher probability of developing polyps (p < 0.01); and older patients showed significantly lower proba-
ibility of polyps in this study (p < 0.001). Further, these results were not affected by life-style related oxidative stress such as smoking and alcohol. Selected SNPs for ox-
idative damage, detoxification and DNA repair were examined. Initial results sug-
ggested a correlation between hOGG1 and GSTM1 and colorectal cancer. The results of this study suggest that high level of oxidative stress induced DNA damage is associated with increased risk of colorectal cancer in males, not in fe-
males.
the occurrence of drug-induced medial arterial necrosis (MAN) in rats; however, a non-invasive method for detecting this type of injury is currently lacking. To determine whether these genes regulated in mesenteric arteries have the potential to be utilized as circulating serum genomic biomarkers, the gene panel was assessed in mRNA isolated from the serum of rats given a vasodose dose of Fenoldopam or Dopamine and compared to mRNA data obtained from serum of rats given Yohimbine, a vasoactive compound which does not cause histologic evidence of vascular injury. Out of the 69 potential circulating biomarkers, 10 genes, including Lrrc59, Tubb6, Kpna2, Abca8a, Pros1, S100a11, IL-8R, CTGF, Nkbfiz and Daf1, were observed to be upregulated (5 fold or greater) and correlated with histologic evidence of MAN following treatment with a vasodose of Fenoldopam. Six of the 10 genes including Lrrc59, Abca8a, S100a11, IL-8R, Nkbfiz and Daf1, were also upregulated (4 fold or greater), in the serum of rats given Dopamine. With exception of S100a11, these genes were not regulated in the serum of rats given Yohimbine thereby, providing further evidence of their association with vascular injury, namely mesenteric MAN. Although further studies are required to fully assess the utility of these serum-derived mRNAs as biomarkers of DIVI, they have the potential to help improve the characterization of DIVI in rats for early safety assessment in drug development.

937 Identification and Quantitative Evaluation of Novel Circulating Liver-Specific mRNAs in Rats Treated with Various Hepatotoxic Compounds: Validation for Biomarkers of Drug-Induced Liver Injury.

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In our previous report, circulating liver-specific albumin (Alb) and tr-1 microglobulin/bikunin precursor (Amblp) mRNAs have been shown to be potential biomarkers for drug-induced liver injury (DILI). We identified apolipoprotein h (ApoH) and group specific component (Gc) mRNAs as additional biomarkers, and quantified total of four mRNAs in plasma from rats treated with wide variety of hepatotoxicants to validate circulating liver-specific mRNAs as biomarkers for DILI. Bioinformatic and molecular biological analyses revealed the high liver-specificity of ApoH and Gc mRNAs, and increased plasma levels of these mRNAs were confirmed by real-time quantitative RT-PCR in rats treated with thioacetamide (TAA). To examine the characters of the four circulating liver-specific mRNAs, seven hepatotoxicants were administered to rats. The severities of liver injury were variable among individuals and compounds. At 24 hr after single dosing, parallel increases of the four circulating liver-specific mRNAs were noted and the levels correlated with changes in the ALT values and hepatocellular necrosis scores. In addition, all the four mRNAs increased with greater magnitude than the ALT values. Time course analysis within 24 hr after single dosing of TAA showed that the plasma levels of Alb and Gc mRNAs increased remarkably before the ALT elevations and the timing of the increase was different among mRNA species, indicating that circulating liver-specific mRNAs may predict the beginning of liver damage and enable us to know the stages of injury. This validation study clearly demonstrated that the four circulating liver-specific mRNAs would be reliable and useful biomarkers for DILI.


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Drug-induced hepatotoxicity represents a major reason that drugs are recalled post market. Furthermore, it has been estimated that 10% of acute liver failure is due to idiosyncratic events. While there is no standard definition for “idiosyncratic,” the term is generally applied to compounds that induce a relatively low incidence of hepatoxicity in humans and fail to exert liver damage using classical toxicity endpoints in commonly used preclinical testing species such as rats and dogs. The inability to identify such compounds with classical preclinical markers of hepatotoxicity has necessitated the need to discover new biomarkers. In order to identify biomarkers of idiosyncratic toxicity, a systems biology study was initiated to evaluate omics endpoints in urine, blood and liver tissue from rats dosed with compounds that have been shown to be overt hepatotoxicans, idiosyncratic hepatotocicants, and negative hepatotoxicants. The two overt hepatotoxicants were acetaminophen (APAP) and carbon tetrachloride (CCL4). Three additional compounds have been studied; one is generally classified as idiosyncratic in nature, ketorolac (FEL), while the other two are considered to not cause liver injury, meloxicam (MEX) and penicillin (PEN). Since idiosyncratic and non-hepatotoxic drugs do not cause overt hepatotoxicity, doses were used to induce some adverse effect (e.g., a decrease in body weight) to provide a phenotypic anchor. Early results show that increases in blood levels of multiple acyl carnitines could be an indication of hepatic mitochondrial injury and altered levels of bile acids may be related to drug-induced hepatotoxicity, activation of liver transporters or due to effects on gut microflora. Blood levels of lysPCs were decreased in the rats treated with a high dose of APAP, CCL4 and FEB at 6 and/or 24 hr.

939 Whole Transcriptome RNA-Seq of FFPE Liver.


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Formalin fixed, paraffin embedded (FFPE) pathology specimens represent a potentially rich resource for transcriptomic-based biomarker discovery. While a number of approaches have been developed that employed targeted, amplification dependent microarray analysis, no one has employed the agnostic approach of whole transcriptome RNA-seq to the analysis of RNA extracted from FFPE samples. Here, we compare whole transcriptome using RNA-seq performed on fresh frozen (FF) and FFPE RNA samples obtained from Fischer F344 rats exposed to Aflatoxin B1 (AFB1) at 1 ppm in feed for 90d and paired control animals. ~70% of reads generated from each FFPE sample aligned to RNA seq and mapping corresponded well with defined transcription boundaries for most RefSeq genes. Whole transcriptome PCA indicated clear distinction between FFPE and FF however robust, parallel differentiation between AFB1 and control samples was obvious in both data sets. A rank based comparison of global transcript abundance was strongly correlated between FF and FFPE, suggesting the global liver biology at the transcript level remained intact in the FFPE samples. Differential expression analysis indicated that of the 405 RefSeq genes altered by AFB1 in FF samples, 288 (of 487 showing differential expression) were also differentially expressed in FFPE samples. Expression of selected genotoxic carcinogenicity biomarker genes (e.g., Adam8, MybH2, Cdh13, Ccn1, Ddr4, Dld1) were concordantly up-regulated in FF and FFPE by AFB1 treatment. Comparison of the genomic response to AFB1 in FF and FFPE at a pathway level using GSEA found concordant regulation of cell cycle, autophagy, xenobiotic metabolism, and p53 signaling and discordant regulation of proteasome and NOD-like signaling pathways. We have demonstrated that accurate whole transcriptome profiling is possible with RNA extracted from FFPE samples. The success of this approach opens up new avenues for performing signature-based biomarker discovery.

940 Hepatic Hemangiosarcoma Due to Occupational Vinyl Chloride Exposure Generates a Distinct Plasma Metabolite Profile.

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Occupational vinyl chloride (VC) exposure has been associated with the development of hepatic hemangiosarcoma (HS), an extremely aggressive and otherwise unusual form of liver cancer. Routine liver biochemistries are typically normal even in advanced hemangiosarcoma. The purpose of this study is to determine if VC-related hemangiosarcoma alters the plasma metabolome. Plasma samples from 16 highly-exposed VC workers with hemangiosarcoma (VC+/HS+), 17 highly-exposed VC workers without hemangiosarcoma (VC+/HS-), and 27 unexposed healthy volunteers (VC-/HS-) were obtained from a specimen bank. GC/MS and LC/MS were performed following metabolite extraction. Software was used to match ions to a library of standards for metabolite identification and quantitation by peak area integration. Random Forest analysis was performed. Welch’s Two Sample t-test comparisons were made between the means of each biochemical. Results of Random Forest analyses of all 613 named and 518 unnamed biochemicals observed in plasma yielded an overall predictive accuracy of 82% for classifying
samples within each group, strongly indicating that hemangiosarcoma generates a distinct global metabolic profile in plasma from those that were exposed but failed to develop the disease. When comparing VC/HS+ vs. VC/HS-, 65 named biochemicals were up-regulated while 50 were down-regulated. Likewise, 43 unnamed biochemicals were up-regulated and 27 were down-regulated. Four key metabolite subclasses containing named biochemical were identified. Hemangiosarcoma increased metabolite levels in these classes accordingly: Class A (8/10 metabolites, up to 6 fold increase); Class B (5/7 metabolites, up to 121 fold increase); Class C (3/5 metabolites, up to 84 fold increase); Class D (9/15 metabolites, up to 92 fold increase). In chemical workers with high-level vinyl chloride exposures, hemangiosarcoma generates a distinct plasma global metabolic profile from those that were exposed but failed to develop the disease.

941 Metabolomic Biomarkers in Long-Term Smokers and Moist Snuff Consumers.
G. L. Prasad1, B. A. Jones1, P. Chen1 and A. D. Kennedy2, 1R & D, RJ Reynolds Tobacco Co., Winston-Salem, NC; 2Metabolon Inc, Durham, NC. Sponsor: G. Krautter.
The long-term health effects associated with cigarette smoking have been shown to be more harmful compared to those associated with the use of non-cancerous tobacco products, such as moist snuff. To investigate the long-term effects of tobacco exposure, we evaluated the biochemical profiles of 40 smokers, 40 moist snuff consumers (MSC), and 40 non-tobacco consumers (NTC) using UHPLC-mass-spectrometry based global metabolomics. Matching twenty-four-hour urine and plasma samples were collected from study subjects to generate metabolomic profiles. In this global profiling study, a total of 511 biochemicals (290 known and 221 unknown metabolites) were detected in the plasma, whereas 972 biochemicals (396 known and 596 unknown) were found in urine. The “named” metabolites could be grouped into diverse physiological pathways such as glucose, lipid, bile acid and amino acid metabolism. Based on the differential levels of metabolites, random forest analyses separated non-tobacco consumers (NTC), smokers, and MSC with high accuracy (96%) when all metabolites were included. On the other hand, MSC showed more subtle changes in metabolomic profiles, and were more difficult to separate from NTC when nicotine metabolites were excluded from the analyses. The metabolic changes suggest that smokers exhibited exacerbated oxidative stress and inflammatory pathways relative to MSC and NTC cohorts. Biochemical changes in glucose, lipid amino acid and xenobiotic metabolism were also noted in the study cohorts.

942 Effect of Benzo(a)pyrene Exposure Dose on Levels of Exposure Biomarkers, DNA Adducts, and Gene Expression in Rats.
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The effect of benzo(a)pyrene (BaP) doses on levels of several biomarkers of exposure and early effects was studied in rats intravenously injected with 0.4, 4, 10 and 40 μmol/kg of BaP. Blood, tissues and excræta were collected 8 h and 24 h post-treatment. BaP and several of its metabolites, 3- and 7-OHBaP, 4,5- and 7,8-BaPdiols, tetrox, 1,6-, 3,6- and 7,8-BaP-diones, were simultaneously measured in blood, tissues and excræta by UHPLC/Fourier Transform IRMS. BaPDE-DNA adducts in livers were quantified in parallel using an ultrasensitive immunoassay with chemiluminescence detection. Expression of various genes in livers of treated rats (lung RNA) compared to control rats was also assessed by QRT-PCR. There was a dose-dependent increase in blood and tissue levels as well as ecretation of BaP metabolites. At 8 h and 24 h postinjection, BaP and 3-OHBaP were found in higher concentrations in blood and tissues compared to the other analytes. However, BaPdiols were excreted in greater amounts in urine and apparently more quickly than hydroxyBaP. Mean percentages (± SD) of injected dose excreted in urine as 4,5-diol-BaP during the 0-8 and 0-24-h period post-treatment were 0.16 ± 0.03% and 0.14 ± 0.08%, respectively. Corresponding values for 3-OHBaP were 0.004 ± 0.001% and 0.026 ± 0.014%. Diones were not detectable in blood, tissues and excræta using the developed method and BaP/7,8-diol and 7-OHBaP were found to be more minor metabolites. There was also a dose-dependent decrease in DNA adduct formation. Analysis of gene expression further showed a modulation of Cyr1a1, cyr1b1, nqo1, nrf2, fos and Ah receptor expression at the 10 and 40 μmol/kg doses, but not at the lower doses. This study confirms the interest of measuring multiple metabolites in combination with DNA adducts and alteration of gene expression for a more comprehensive assessment of links between biomarkers of BaP exposure and early effects.

943 Biomarker Discovery for Early Detection of Hepatocellular Carcinoma (HCC) in Hepatitis C (HCV) Infected Patients.
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The projected rise in HCC (the most common primary liver cancer) cases in the US is mainly due to HCV infections with onset of HCC coming several decades after initial infection. However, additional environmental risk factors including alcohol, tobacco and other dietary insults that induce liver injury also increase the incidence of HCC. The poor prognosis for HCC is largely due to late stage diagnosis making successful intervention difficult. Existing biomarkers for early HCC detection lack the specificity and sensitivity to be very effective. Our aim is to develop serum-based biomarkers suitable for early HCC detection that will provide a sensitive yet specific screening. Serum was obtained from individuals positive for HCV who were clinically diagnosed with liver disease (pre-HCC) or HCC. Serum was pre-fractionated using an amptzer-based technology and further fractionated using 2D-Difference In Gel Electrophoresis. Pre-HCC and HCC profiles were compared and the peptides located in 24 different 2D-DIGE spots exhibiting a statistically significant ≥1.5-fold change between pre-HCC and HCC samples were identified by mass spectrometry (MS). Stable isotope O18/O16 labeling was used to verify the identity of proteins identified by 2D-DIGE and to aid in development of Selected Reaction Monitoring (SRM) assays. ApoA1 was selected to develop an SRM as proof of concept in this biomarker discovery protocol. Using a Triple Quadrupole MS (Agilent), Optimizer (Agilent) and skyline (MacCos) to assist in the design we developed an SRM assay to quantify this candidate biomarker using labeled internal standards (AQA peptides). The SRM is capable of reliably detecting a >30% reduction in ApoA1 in HCC serum samples relative to the pre-HCC samples.

944 Effects of Acrylamide Exposure on Gene Expression in the Thyroid of Male and Female Rhesus Wistar Rats.
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Acrylamide is a chemical commonly utilized to make polymers used in industry including waste water treatment, mining, among other uses. It became a public health concern when it was detected in food products. Daily exposure to acrylamide has been associated with neurological toxicity in different animal species and with tumors in the mammary gland, testicular tumica vaginalis, and thyroid. In relation to potential for forming tumors, it is important to distinguish its ability to act as a genotoxin from a hormone mimic. The goal of this study was to use microarray analysis to determine changes in gene expression in the thyroid gland of male and female RecHan Wistar rats treated at 3 mg/kg in drinking water from gestational day 6 to postnatal day 30. Thyroid glands were collected from rats at 10 PM to reflect end of non-quinsecent activity. The transcriptome analysis showed that acrylamide caused a significant alteration in the expression of genes related to phase II detoxifying enzymes and oxidative stress, neurotoxicity, apoptosis, and tumorigenesis. Results from enrichment and pathway analysis showed that in both genders, acrylamide caused differential regulation of genes associated with cell processes of protein folding, microbule cytoskeleton assembly, DNA replication, DNA repair and ROS generation. These data suggest a potential involvement of this chemical in cell processes other than mutagenicity which may be associated with tumorigenesis. Furthermore, our study, by identifying affected gene ontologies and pathways, provides insight into a potential mechanism of action and may have substantial impact on risk assessment.

945 Development of a Quantitative Proteomics Multiplexed Assay for the Predictive Assessment of Drug-Induced Organ Toxicity in the Preclinical Setting.
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Drug-induced organ toxicity is the primary reason for the withdrawal of drugs from the market and the failure of lead compounds during drug development. Detection of organ toxicity early in the development pipeline would not only reduce the cost
of drug development but also prevent injuries to patients in the clinic. Limitations of traditional approaches highlight the need for more sensitive and specific tools to predict drug toxicity. With the recent regulatory qualification of seven renal safety biomarker for use during drug development, biomarkers are increasingly viewed as potential means for providing toxicity assessments. Here, we have employed a targeted mass spectrometry (MS) approach, known as multiple reaction monitoring (MRM) to quantitatively measure changes in organ-specific toxicity biomarkers in rats. A highly multiplexed MRM assay, targeting 192 candidate protein biomarkers of hepatotoxicity, nephrotoxicity, cardiotoxicity as well as muscle, vascular and neurotoxicity, was developed. Results show that markers for all organs covered in our toxicity biomarker panel are detectable in serum and urine under normal conditions. Significantly, Cc105 and MDH-1, important liver injury biomarkers currently under investigation as well as qualified kidney injury biomarkers cystatin-C, clusterin and β-2-microglobulin and albumin were successfully detected in serum and urine, respectively. We tested the predictive value of our panel on cyclophosphamide treated rat blood samples. In agreement with the literature, we found that serum biomarkers for the kidney, the vascular system and the liver were markedly increased following toxicity. The present MRM assay has the potential to become standard practice for pre-clinical drug safety assessments. Further studies will be centered on applying this assay to monitor urinary kidney injury biomarkers following cisplatin-induced nephrotoxicity of rats.

946 Development of a LC/MS Serum Bile Acid Profiling Method for Sensitive Detection of Biliary Injury in Dogs.


Currently there are many non-invasive methods for assessing hepatic toxicity in large animal species, but biliary injury is often difficult to detect in lieu of histopathology. Notably hepatobiliary biomarkers such as liver transaminases, alkaline phosphatase, gamma glutamyl transferase, and total bilirubin suffer from poor specificity or sensitivity in detection of biliary injury. Quantification of serum bile acids has been utilized in veterinary medicine in the diagnosis of biliary disease with success, but the current methods of quantification suffer from technical hurdles and lack robustness. Thus the purpose of the current study was to develop a method for detection of serum bile acids in dogs using liquid chromatography-mass spectrometry (LC/MS). We performed rapid detection of hepatobiliary injury by quantifying the individual bile acid species in tandem. Twenty bile acids species were assessed with the current LC/MS method, including primary cholic and chenodeoxycholic bile acids and secondary bile deoxycholic and lithocholic bile acids, including the glycine- and taurine-conjugated forms. A diagnostic reference range was established from more than 50 untreated, fasted beagles of both genders to assess normal variability with benchmarked clinical pathology. These individual serum bile acid levels ranged from approximately 900 pmol/ml to levels below the limits of detection, with all taurine-conjugated species notably detected in serum. The applications of this method in assessing the sensitivity and specificity of serum bile acid versus existing methods will be discussed in the early detection of drug-induced hepatobiliary injury.

947 Lack of Concordance in Microarray Gene Expression Responses to Phenobarbital in Companion-Aged FFPE and Frozen Liver Samples.

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Despite the immense potential value of public and private biorepositories, direct utilization of archival tissues for molecular profiling has been limited. A major reason for this limited use is the difficulty in obtaining reliable transcriptomic profiles from formalin-fixed paraffin-embedded (FFPE) tissue samples with highly fragmented nucleic acid. The goal of this study was to evaluate transcriptional responses in 16 year-old FFPE and companion frozen (FROZ) liver samples from F344 rats treated for 2 yrs with control water or water with 0.06% Phenobarbital (PB), a known CAR/PXR inducer and rodent liver mitogen. RNA isolation and labeling methods were optimized for FFPE tissues were used to amplify and label the RNA followed by hybridization to arrays. We selected 12 of 16 samples based on successful hybridization performance for analysis. Differential gene expression was assessed using Rank Products followed by pathway mapping. Results showed only 10 significantly altered PB-responsive genes in common between paired FFPE (n=1373) and FROZ (n=21) samples whereas pathway analysis identified only a 10% overlap in pathways significantly altered by PB between FFPE and FROZ samples. Pathway profiles for both FFPE and FROZ samples were unique to their respective sample type. The lack of concordance in genomic responses to PB suggests that traditional microarray platforms are inadequate for genomic profiling of aged FFPE samples, despite improved RNA isolation and labeling methods. Further work will evaluate more recent next-generation sequencing (NGS) technologies for transcriptomic analysis of archival specimens. This abstract does not reflect EPA policy.

948 Formaldehyde-Induced Changes in microRNA Signaling.

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MicroRNA (miRNA) are critical regulators of gene expression, yet much remains unknown regarding miRNA changes resulting from environmental exposures and whether they influence pathway signaling across various cell types and tissue. To gain knowledge on these novel topics, we set out to investigate in vivo miRNA responses to inhaled formaldehyde, an important air pollutant known to disrupt miRNA expression profiles. Rats were exposed by inhalation to either 0 or 2 ppm formaldehyde (6 hours/day) for 7 days, 28 days, or 28 days followed by a 7 day recovery. Genome-wide miRNA expression profiles and associated signaling pathways were assessed within the nasal respiratory mucosa, circulating mononuclear white blood cells (WBC), and bone marrow (BM). We found that miRNAs were responsive to formaldehyde exposure in the nose and WBC, but not the BM. A transcriptomics-based analysis was performed in the nose and WBC of the rats exposed for 28 days. In the nose, formaldehyde altered the expression of 42 transcripts; of these, 15 (36%) were computationally predicted to be regulated by formaldehyde-responsive miRNAs. Conversely, in the WBC, formaldehyde altered the expression of 130 transcripts; of these, 18 (14%) were predicted to be regulated by miRNAs. Systems-level analyses revealed that the transcripts regulated by miRNAs play diverse roles in cell signaling. Key players include dosage suppressor of mck1 homolog, meiosis-specific homologous recombination (Dmc1) and secreted frizzled-related protein 4 (Sfrp4) within the nose, involved in cell death signaling. In WBC, key players were v-akt murine thymoma viral oncogene homolog 3 (protein kinase B, gamma) (Akt3) and integrin, alpha 2 (Itga2), involved in inflammation signaling. Our study informs critical knowledge towards the biological consequences of inhaled formaldehyde exposure.

949 Dermal Sensitization to Toluene Diisocyanate in Mice Is Manifested by Early Changes in the Proteome of Auricular Lymph Nodes and Serum.

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Introduction.
Diacyanates are capable of initiating an allergic response, which can lead to occupational asthma after a latency period. Clinical symptoms such as cough, wheezing and dyspnea occur only later, making it difficult to intervene at an early stage. We explored proteome changes, before asthma is apparent, in the local draining lymph nodes and serum of mice dermally sensitized once or twice with toluene diisocyanate (TDI) to explore biomarkers of sensitization.

Methods.
The proteomes of male BALB/c mice (6 weeks old, 20g, sensitized once (n=12) or twice (n=12) with 0.3% TDI, were individually compared with control mice (n=12) using two-dimensional difference gel electrophoresis (2D-DIGE). A level of p<0.05 (Student’s t-test) was considered significant.

Western blot or ELISA was used to verify a subset of the differential proteins. A level of p<0.05 (unpaired t-test) was considered significant.

Results.
In the lymph nodes, we found between TDI-treated and control mice 38 and 58 differential proteins after one and two treatments, respectively. In serum, 7 and 16 differential proteins were detected after one and two treatments, respectively. We identified 80-85% of these proteins by mass spectrometry. Among them, lymphocyte specific protein-1, coronin 1a and hemopexin were verified in an independent group of mice by Western blot or ELISA.

Conclusion.
Our study revealed, in a mouse model, alterations in the proteome during sensitization. If validated in humans, these changes could lead to earlier diagnosis of exposed workers. All animal experiments were approved by the Local Ethical Committee for animal experiments of the KU Leuven.
950 Smoking-Induced microRNA Changes in Human Sperm.  
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Recent work has suggested that some of the constituents of cigarette smoke, along with other environmental chemicals, can have adverse effects not just on the exposed individuals, but also on their progeny. Although the mechanisms underlying multigenerational toxicity are not well understood, a number of studies have implicated heritable microRNA-mediated epigenetic modifications. Using microarray profiling and pathway analysis, we have shown that cigarette smoke induces specific differences in the spermatozoa microRNA content of human smokers compared with non-smokers, and that these altered microRNAs appear to predominantly mediate pathways vital for healthy sperm and normal embryo development, particularly cell death and apoptosis. MicroRNA-mediated perturbation of such pathways may explain how harmful phenotypes can be induced in the progeny of smokers. Consequently, we have also been developing an in vivo system for investigating the potential roles of microRNAs in toxicology. By differentiating embryonic stem cells into embryoid bodies we have been able to generate and subsequently isolate sets of different embryoid bodies we have been able to generate and subsequently isolate sets of clones representing a rich and largely untapped resource of invaluable information. There is a need to develop methods to study the effects of smoking on the spermatozoa microRNA content and elucidate the potential adverse effects of smoking on sperm and offspring. A more comprehensive understanding of the potential for smoking-induced microRNA changes and the mechanisms underlying such changes may lead to the development of novel biomarkers for smoking-induced adverse effects on reproductive health.

953 Determination of Formaldehyde Specific DNA-Protein Crosslinks.  
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Formaldehyde (FA) is a known human and animal carcinogen. It is a ubiquitous environmental pollutant and is used in a number of consumer products or industrial applications. FA is also endogenously produced as part of normal cellular metabolism. FA is a genotoxic agent, causing a number of effects on cells including DNA monoaducts, DNA-DNA crosslinks and DNA-protein-crosslinks (DPC). DPCs are believed to be one of the critical lesions involved in FA induced carcinogenesis and is thought to provide a key initiating step in the Mode of Action. Currently, there are no available methods to distinguish between endogenous and exogenous FA induced DPCs. To investigate the possible link between endogenous FA and the formation of both endogenous and exogenous DPCs, several analytical techniques are being developed. O-Alkylguanine-DNA alkytransferase (AGT) is a DNA repair enzyme that is known to form DPCs with FA and other crosslinking agents at the active Cys145 residue. Using AGT as a model protein, a series of experiments were undertaken to understand the formation, stability and degradation of the DNA-protein-crosslink at the reactive cysteine and the N2 position of deoxyguanosine. Digestion conditions for both DNA and protein cleavage were investigated to determine approaches that would allow for the isolation and identification of either cysteine–CH2–dG or AGT peptide–CH2–dG crosslinks using sensitive and selective Liquid Chromatography – Mass Spectrometry. Further experiments investigating the ability to distinguish between crosslinks formed by both [13CD2]-FA and unlabeled FA were accomplished. Further validation and development of these approaches may allow for accurate and quantitative determination of endogenous and exogenous FA specific DPCs in cell culture and animal models. This information will be critical in advancing the understanding of the risks associated with inhaled FA and its role as a human carcinogen.

954 Pulmonary Toxicity and Global Gene Expression Profile in Response to Crystalline Silica Exposure in Rats.  

The ability to detect target organ toxicity as well as to determine the molecular mechanisms underlying such toxicity by employing surrogate biospecimens that can be obtained either by a non-invasive or minimally invasive procedure has significant advantage in toxicology. Pulmonary toxicity and global gene expression profiles in the lungs, blood and bronchoalveolar lavage (BAL) cells were determined in rats 44-weeks following inhalation exposure to crystalline silica (15 mg/m3, 6-hours/day, 5 days). A significant elevation in lactate dehydrogenase activity and albumin content in the BAL fluid as well as histological alterations, mainly type II pneumocyte hyperplasia and fibrosis, observed in the lungs suggested silica-induced pulmonary toxicity in the rats. A significant increase in the number of neutrophils and elevated monocyte chemotactic protein 1 in the BAL fluid indicated silica-induced pulmonary inflammation in the rats. Determination of global gene expression profiles in the lungs, BAL cells, and blood of the silica exposed rats identified 175, 273, and 59 significantly differentially expressed genes (SDEGs) (FDR p<0.05 and >1.5 fold change in expression), respectively, compared with the corre-
An essential step in the safety review of cosmetic/personal care ingredients is hazard assessment, including dermal sensitization potential. In vitro methods to identify receptor fluid.

Critical endpoints in in vitro testing of cosmetic ingredients are the determination of the bioavailability of test substances in different skin layers and the examination of the toxicokinetic profile. Skin penetration studies are so far performed in Franz diffusion cells using pig skin. Unfortunately with these cells an automated toxicokinetic determination of percutaneously penetrated substance is not receivable.

To perform toxicokinetic studies, we developed a new Vitrocell systems skin penetration system (SPS) with eight parallel running diffusion cells, which is able to take samples from the receptor fluid automatically. To substitute the Franz diffusion cells with the SPS it is important to compare both systems in terms of performance and reproducibility. Therefore we compared the penetration of caffeine through full thickness pig skin in Franz diffusion cells with manual sampling from the receptor fluid with the prototype of the SPS that provides automated sampling from the receptor fluid.

We could show toxicokinetic profiles for manual and automated samples with comparable lag times and recovery rates. Furthermore we could even show lower standard deviations using the SPS.

In conclusion, the new SPS is highly comparable to the Franz diffusion cell with the additional advantage to allow the automated detection of toxicokinetic profiles from the receptor fluid.

Development of non-animal safety evaluation methods for chemicals is necessary from the viewpoint of animal welfare and to meet the 7th amendment of the European cosmetic directive. As an in vitro skin sensitization test, we have established two methods. One was the human cell line activation test (h-CLAT) detecting augmentation of CD86 and CD54 expression in THP-1 cells exposed by skin sensitizers. Recently the Antioxidant Response Element (ARE) assay measuring oxidative stress caused by skin sensitizers have been described two methods. One was the human cell line activation test (h-CLAT) detecting augmentation of CD86 and CD54 expression in THP-1 cells exposed by skin sensitizers. Recently the Antioxidant Response Element (ARE) assay measuring oxidative stress caused by skin sensitizers have been described. The second method was selected because it operates over a wide dose range and sets cytotoxicity limits which is currently validated for classification of chemicals for dermal irritation in OECD 439, and also dermal corrosion in OECD 431. Following the process outlined in the standard protocols for demonstration of technical proficiency, both test methods have been validated and used to assist with classification of chemistries for consumer and industrial use, utilizing the standardized MTT based cell viability assay. In addition, custom protocols have been performed using the tissues in time course irritation studies to evaluate the release of the inflammatory marker cytokine.

Application of a Modified Keratinosens Assay to Predict Sensitization Hazard for Botanical Extracts.

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An essential step in the safety review of cosmetic/personal care ingredients is hazard assessment, including dermal sensitization potential. In vitro methods to identify allergic (haptenic) potential are based on electrophilic interaction with marker peptides or cellular target systems. These assays use a specific molar ratio of the test chemical to the test system such as mixtures, which preclude specific molar ratio determination. Often, the botanical extract portion is a small fraction of the complete ingredient. To assess these mixtures, the Keratinosens assay was selected because it operates over a wide dose range and sets cytotoxicity limits on doses used to measure marker gene expression (Emter et al, 2010). Induction of a downstream gene, under the control of the antioxidant response element (ARE), was measured. Cytotoxicity was assessed by both NRU and MTT assays. Concentrations up to 1 mg/mL (of complete ingredient) were tested and a test dose was considered positive if the fold induction of luciferase was 1.5x (EC1.5) and viability ≥ 70% relative to the solvent controls. The goal of the study was to measure the activity of 3 known sensitizers (gluteraldehyde [GA] strong), dimethyl maleate [DM] [moderate] and cinnamon aldehyde [CA] [moderate] spiked into four different botanical ingredients (with different excipient solvent systems). Activity was measured, relative to the EC1.5 of the neat sensitizer, as a function of sensitization concentration and ingredient composition. Three independent trials were performed. No appreciable cytotoxicity was observed. The recovery of the GA spike required at least a 3-fold increase in concentration relative to the chemical alone and one extract reduced the activity below detection. The DM and CA showed activity at about the same effective concentrations as the neat chemicals although the DM showed reduced activity in one extract as well. These data suggest that the Keratinosens assay has the potential to identify electrophile allergens within a botanical extract ingredient matrix.

In vitro models to study irritation, corrosivity and phototoxicity are important tools for research and development in the pharmaceuticals and cosmetic industries. Human skin is the best possible model for such in vitro studies. The commercially reconstructed human epidermis models are similar to the morphology, lipid composition and biomarkers of native human tissue and have been approved by European Centre for the Validation of Alternative Methods (ECVAM) for the validation of cosmetics. The models are available in many countries but not in China. Here, we describe the development of a constructed three-dimensional (3D) model using Chinese human skin, which consists of a ‘dermis’ with fibrobloasts embedded in rat collagen matrix and an ‘epidermis’ comprised of differentiated keratinocytes. The fibrobloasts and keratinocytes were first separated from foreskins of Chinese adults after incubating in dispase and collagenase solutions. Then, rat type I collagen was constructed onto the polycarbonate membrane of a culture insert. After gels solidified at room temperature, a collagen matrix with Chinese dermal fibroblasts was constructed above the acellular collagen layer. Keratinocytes were added to the surface of the collagen matrix in alginate capsules to attach to the matrix to create a coherent cellular monolayer. The reconstructed tissues were irrigated with an air-liquid interface to enable complete stratification and differentiation. After exposure and incubation, MTT assay was performed for cell viability. Nine well-known irritants or corrosive chemicals caused cell viability rates less than 50%. Cosmetics both from Western nations and China were also tested in the model. We found the Wafa North herbal cream did not show toxicity or viability up to 0.5% of some of Chinese cosmetics which dramatically decreased cell viability 1 hour after exposure. Our data indicate that the reconstructed 3D Chinese skin model is highly reproducible and sensitive to assess skin irritation to chemicals and cosmetics.

Incorporation of Reconstructed Human Epidermal Tissues into a Corporate Toxicology Laboratory: Use of In Vitro Test Data for Diverse Safety and Risk Assessment Applications.

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The 3M Strategic Toxicology Laboratory (STL) is an internal corporate resource providing toxicology support to 3M businesses. An objective in the STL is to routinely incorporate in vitro test methods, with a recent focus on the use of reconstructed human epidermis tissues. The first model utilized has been Epiderm™, which is currently validated for classification of chemicals for dermal irritation in OECD 439, and also dermal corrosion in OECD 431. Following the process outlined in the standard protocols for demonstration of technical proficiency, both test methods have been validated and used to assist with classification of chemistries for consumer and industrial use, utilizing the standardized MTT based cell viability assay. In addition, custom protocols have been performed using the tissues in time course irritation studies to evaluate the release of the inflammatory marker cytokine.
IL-1α. This approach has been advantageous for the simultaneous screening of multiple formulations for irritation potential during research and development for new products. Other customized studies have evaluated the efficacy of various skin washing agents at removal of a viscous coating from the skin tissues, in a simulated washing study to help formulate occupational health recommendations for workers. These studies evaluated irritation potential using MTT, IL-1α and LDH release, and also analyzed the dermal penetration of an amine component in the coating. In summary, incorporation of these test methods has been very useful for the evaluation of a multitude of dermal exposure scenarios. The tissues have been used in standardized irritation and corrosion assays, as well as custom protocols utilizing additional markers of cell damage, and also specialized skin washing and dermal penetration studies. The results of these studies have provided valuable information for safety and risk assessment purposes, particularly during the product development phases, and have permitted these evaluations without animal use.

**960 Intralaboratory Reproducibility of the Direct Peptide Reactivity Assay (DPRA) to Assess the Robustness of the Current Prediction Model.**

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One characteristic of a chemical allergen is its ability to react with proteins prior to the induction of skin sensitization. The majority of chemical allergens are electrophilic and react with nucleophilic amino acids such as cysteine or lysine. In the DPRA, test chemicals are incubated for 24 hours with two synthetic peptides containing a cysteine or lysine residue, and the reactions are analyzed by HPLC to measure peptide depletion. We have previously demonstrated that reactivity correlates with sensitization potential and have developed a prediction model to assign a reactivity category for each material tested in the assay (Gerberick et al. *Tox Sci* 2007; 97:417-427). In order to assess the robustness of this prediction model and its ability to distinguish reactive skin sensitizers from minimally reactive non-sensitizers, a historical DPRA dataset of 133 chemicals was analyzed retrospectively with a modified prediction in which the cut-off between minimal and low reactivity was changed from 630% to 8% depletion. Regardless of cut-off, the accuracy of the prediction model remained at 86% with little to no change in sensitivity, specificity, and positive and negative productivity. To further investigate the robustness and reproducibility of the prediction model, a small intra-laboratory study was conducted. In this study, 9 chemicals were tested blindly in three independent runs. The depletion data was analyzed using the current prediction model as well as the described modified prediction model. In both cases, 7 out of 9 chemicals showed the same reactivity prediction in all 3 runs. Neither the retrospective analysis nor the intra-laboratory study provided any compelling evidence to adjust the cut-off value from the current model. Taken together, these studies further display the reproducibility and robustness of the DPRA and its prediction model.

**961 A Novel Animal-Free Approach to Dermal Sensitization Hazard Identification and Potency Assessment.**


International regulatory efforts have played a major role in advocating for the Three Rs of animals used in safety evaluations for toxicological endpoints. These efforts support a movement away from the current in vivo-dependent toxicity testing state, to an animal-free toxicity testing state that includes in vitro human biology and a computational systems approach. To accomplish this goal, key stakeholders have proposed an ITS methodology to assess the skin-sensitizing potential of test chemicals to facilitate safety evaluation decision-making. The use of this ITS model integrated both novel and literature-reported data on 116 chemicals of known skin-sensitizing potential. Two in vitro assays and one in silico prediction model were used to cover four key sources of toxicological information: peptide reactivity; induction of ARE dependent luciferase activity; catalysis of transcutaneous water partition coefficient; and TIMES-SS model in silico prediction. The results from the in vitro assays were scaled into five classes from 0 to 4 and then compared with the LLNA data, which were also scaled from 0 to 4. Using an optimized model, an 87.9% overall accuracy was obtained for predicting sensitizers with a linear relationship between the LLNA scores and the in vitro scores. The use of clogP to account for potential differences in the bioavailability of the test chemicals did not reduce the correlation. With the increasing number of large-scale toxicology risk assessment programs for chemicals and emerging or expanding areas of chemical use, the development and acceptance of an ITS model appears to be the solution needed to strike a balance between safety, speed, cost and animal welfare. Along with ITS data refinements and the inclusion of other in vitro assays and in silico models accompanied by larger data sets, consideration should be given to defining a validation strategy focused on regulatory acceptance of an ITS model for skin sensitization.

**962 Development of a Compromised Skin Model In Vitro Using a Tape Stripping Method.**

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Dermal absorption of chemicals from topically applied products is studied in a variety of in vitro and in vivo models. Skin barrier function may be impaired due to intrinsic or extrinsic factors. To our knowledge there is not a standardized model for evaluation of dermal absorption in compromised skin barrier conditions. Our objective was to validate an in vitro model of altered barrier function generalizable across compromised skin states. To investigate the permeability properties of compromised skin compared to normal skin, we explored the relationship between non-invasive (Trans Epidermal Water Loss, TEWL) and invasive (Electrical Resistance, ER; Tritiated Water Flux, TWF) markers of barrier function in slaughterhouse pig skin before and after tape stripping. Skin samples were randomly divided into groups and subjected to a standardized tape stripping procedure. The pre-and post-values for the 3 measures of skin integrity were recorded for 0, 5, 10, 15 and 20 tape strips. A full analysis of the distribution of TEWL, ER and TWF for 70 separate skin samples revealed that only ER was robust enough to discriminate between the barrier property changes effected by sequential tape stripping. The measurement of water flux through the skin (TEWL, TWF) required a long period of stabilization and proved to be unsuitable as a short term test. However, a significant difference could be observed with ER between control and each group of 5-20 tape strips (p<0.01, n = 53). Further analysis of the data revealed that removal of 10 tape strips provided a loss of barrier function approximately equivalent to a 3-4 fold increase in TEWL, which approximates the altered barrier function clinically observed in atopic dermatitis, psoriasis, and diaper dermatitis. In conclusion, we have developed an in vitro model of compromised skin barrier function that is simple and robust and can be used to study the dermal absorption of chemicals that may come in contact with skin with impaired barrier properties.

**963 A Predictive High-Content Imaging Cell-Based Assay as an Alternative to the Local Lymph Node Assay for Skin Sensitization.**


Allergic contact dermatitis (ACD) is a health effect that can develop in those exposed to skin sensitizing chemicals. To decrease the occurrence of this adverse reaction, regulations require that testing be performed to identify which chemicals are responsible for this effect, and be labeled accordingly. The current standard for ACD measurement is the Local Lymph Node Assay (LLNA), which is both animal and labor intensive. The LLNA testing procedure is based upon enumerating the responding cells in the draining lymph nodes of the mouse after treatment. Alternatives for this assay have depended on the measurement of phenotypic changes of a defined cell line, which would be analogous to a prior step in the pathway of induction of sensitization than that of the LLNA endpoint. To the end of developing an assay that more comprehensively mimics the role of the immune system during ACD in vitro, we describe a phenotypic and functional cell based assay that can be used to predict the sensitizing potential of compounds. The prediction method of this assay is based upon the up-regulation of cell surface molecules and proliferation indicative of the induced maturation of the cell line after compound treatment, and also a functional response whereby the treated cells are exposed to chemotaxis inducing reagents and migration measured. These features can easily be captured and measured via an automated fluorescent imaging and analysis platform. The measurement of these features when combined allow for prediction of sensitizing potential, but also provide toxicity data which allows for compounds to be easily re-screened at lower concentrations. Tested compounds (n=24) yielded a high sensitivity rate (100%) and a high concordance (82%) to the LLNA. In summary, this assay shows promise in its ability to predict in vivo responses and may offer a viable alternative to traditional animal studies. In addition, it could provide a valuable first level triage for chemicals entering the consumer and industrial products pipeline.
To encourage the development and validation of alternative toxicity test methods, the effort required for validation of test methods proposed for regulatory purposes should be minimised. Performance standards (PS) facilitate efficient validation by requiring limited testing. Based on the validated method, PS define accuracy and reliability values that must be met by the new similar test method. The OECD adopted internationally harmonized PS for evaluating new endpoint versions of the local lymph node assay (LLNA). However, in the process of evaluating a lymph node cell count alternative, the LNCC, simultaneous conduct of the regulatory LLNA showed that this standard test may not always perform in perfect accord with its own PS. The LNCC results were similar to the concurrent LLNA; discrepancies between PS, LLNA and LNCC were largely associated with “borderline” substances and the variability of both endpoints. Two key lessons were learned: firstly, the understandable focus on substances close to the hazard classification borderline are more likely to emphasise issues of biological variability, which should be taken into account during the evaluation of results; secondly, variability in the results for the standard assay should be considered when selecting reference chemicals for PS.

### Alternative Method in Practice: Postvalidation Experience of the Skin Sensitization In Vitro Test Strategy.

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Several in vitro methods including dendritic cell line activation (e.g. MUSST and h-CLAT), keratinocyte activation (e.g. LuSens and Keratin0Sens) and in chemico (e.g. DPRA) assays have been described as promising animal-free tools to qualitatively predict skin sensitizing potential. While these methods are currently undergoing evaluations in the different stages of formal validation, testing strategies have been proposed based on the combination of these assays. Yet to use suggested methods and prediction models for the diverse industrial sectors, such as the cosmetic, industrial chemical, pharmaceutical and maybe even the agrochemical sector, the scope of the substance classes tested as part of the initial validation exercise needs to be extended. Typically in a first validation phase novel alternative methods are evaluated against their gold standard in vivo assays using model substances selected from literature for their well described toxicological endpoint effects. The substances tested in the first validation phase, do, however, usually not reflect the typical test substance portfolio of the different industrial sectors. Therefore in this study we present the post-validation evaluation of 40 additional substances with available in vivo skin sensitization data from various substance classes including acrylates, surfactants, isocyanates, plant extracts, and agrochemical formulations in state of the art in vitro methods to assess skin sensitization. This additional data provides valuable information to understand the predictive capacity in terms of the applicability domains and may also help to manage expectations what can be achieved with those assays.

### Evaluation of Skin Irritation for Household Products Using a Reconstructed Human Epidermis Model.


Classification for irritation using reconstructed epidermis test methods have been developed. These test methods have been validated with the ability of dividing chemicals into skin irritants or non-irritants, in accordance with the UN Globally Harmonized System of Classification and Labeling (GHS). Application of these models to dermal safety evaluation of household products, which is consisted of several chemical substances, is of highly importance from the point of view to ensure product safety. The aim of this research is to estimate skin irritation potential of household products comparing cell viability patterns on reconstructed epidermis when household products are applied.

We select LabCyte EPI-MODEL24 (Japan Tissue Engineering Co., Ltd) model with JACVAM validated protocol for GHS classification of chemical substances. Twenty five commercial household detergents in Japan including laundry detergents, dishwashing detergents, fabric softeners and household cleaners were tested. It appeared that the cell viability becomes comparable in some product categories. The products which contains highly anionic surfactants tends to show low cell viability. Fabric softeners which were based on cationic surfactants showed no cytotoxicity in this model. Powder laundry detergents tend to show low cytotoxicity. Liquid laundry detergents, dishwashing detergents and household cleaners show various cell viability. It seemed that category and/or its concentrations of contained surfactants might be affected the results.

These results suggested that reconstructed epidermis test methods useful for the products safety assessment by performing comparative assessment in each product category.
For the replacement of the LLNA, the combination of several alternative methods is necessary. JCIA organized a working group to investigate testing strategies. In this study, considering the applicable domain, we demonstrated the utility of a testing strategy using the human Cell Line Activation Test (h-CLAT), Direct Peptide Reactivity Assay (DPRA) and the in silico system, DEREK. A total of 133 chemicals, some of which exhibit poor water-solubilities, were evaluated. For h-CLAT, THP-1 cells were exposed to each test chemical for 24 hours. The CD86 and CD54 expressions were analyzed by flow cytometry. For DPRA, model peptides were mixed with each test chemical for 24 hours. The depletion of peptides was analyzed by HPLC. For DEREK, the alert of chemical structure was examined. For 133 chemicals, the accuracy of h-CLAT, DPRA and DEREK to predict LLNA results was 78, 74 and 74%, respectively. Next, we investigated 3 testing strategies: the Integrated Testing Strategy (A), a tiered approach (B), and a multiple regression analysis model (C). Each strategy had a high potential prediction (A:84%, B:81%, C:84%). Regarding false negative chemicals, some chemicals induced no cytotoxicity, and M. Miyazawa1,11. K. Okamoto11, N. Imaz1,11, D. Kyotani5,11, Y. Kato5,11, T. Kasahara5,11, A. Toyoda5,11, S. Watanabe5,11, Y. Seto5,11, T. Ashikaga5,11 and M. Miyazawa11,11. Kao Co., Tokyo, Japan; 2Kose Co., Tokyo, Japan; 3Cosmos Technical Co., Ltd., Tokyo, Japan; 4Nippon Menard Cosmetic Co., LTD., Aichi, Japan; 5FujiFilm Co., Kanagawa, Japan; 6P&G Japan K.K., Hyogo, Japan; 7Shiseido Co., LTD., Kanagawa, Japan; 8Working Group for In Vitro Skin Sensitization Evaluation in Japan Cosmetic Industry Association (JCIA), Tokyo, Japan. Sponsor: J. Avalos.

By line in the 3Rs concept, in vitro alternatives methods are being developed based on the early events of the skin sensitization process. One of these is the capacity of skin dendritic cells to recognize a chemical as a danger. Since the late nineties, L’Oréal has actively worked on the Myeloid U937 Skin Sensitization Test (MUSST) which models this early event by measuring the up-regulation of CD86 expression, a co-stimulator receptor. Year’s in-house use of this assay allowed us to demonstrate its value in combination with other assays (DPRA, Nf-2 activation) for the prediction of sensitization hazard (Sensitizer S /Non-Sensitizer NS) of cosmetic ingredients. A third class was defined as “Inconclusive (INC)” where the MUSST is not able to conclude S or NS. The aim of this study was to refine the predictive model in order to reduce the number of INC, by maintaining the predictive capacities of the MUSST. 

Methods: 208 substances with LLNA data were analyzed. MUSST results on this set gave 92.5, 56 NS predictions and 60 INC. Statistical methods based on a stacking scoring method were applied on raw data of these 60 INC in order to find additional rules to classify these substances into S or NS. 

Results: a combination of 6 rules was identified, that allows to classify 44/60 INC into 23 S and 21 NS with a sensitivity of 95% and a specificity of 91%. The predictive performances of the MUSST on the 208 substances with the optimized prediction model composed of the original prediction model and the additional 6 rules-based model show 94% concordance, 87% sensitivity and 97% specificity against LLNA data. No false negative are observed among the extreme and strong sensitizers. 

Conclusion: The optimized MUSST is an efficient assay for the skin sensitization hazard characterization, promising as a tool to be integrated within a battery of assays to perform a skin sensitization risk assessment.

According to current regulatory guidelines, photosafety testing is required for new compounds if they absorb light in the range of 290–700 nm or partition into the skin or eyes. The only approved non-animal in vitro phototoxicity assay, the 3T3 Neutral Red Uptake (3T3 NRU-PT), is overly-sensitive in predicting the in vivo photosafety hazard to humans and thereby eliminates valuable, new active pharmaceutical ingredients (API) from further development, even though they are safe to humans. EpiDerm™, a normal human 3-dimensional (NHu-3D) skin model, is highly reproducible, contains an in vivo-like barrier, possesses in vivo-like biotransformation capabilities, and has been pre-validated for determining phototoxicity of topically applied materials. Here, we utilized EpiDerm to develop an in vitro assay for screening the phototoxicity of pharmaceuticals following systemic administration (iPHO). Test articles (n=42) were added into the culture medium and allowed to partition into the epidermal tissue. Tissues were exposed to solar radiation and phototoxic effects were determined by the comparing the tissue viability of UV irradiated vs. non-irradiated tissue models, using the MTT assay. A prediction model (PM) was established: a material is phototoxic after systemic administration if one or more test concentrations in the presence of irradiation (+UVR) decreases tissue viability by ≥30% when compared to identical concentrations in the absence of irradiation (-UVR); a material is non-phototoxic if the decrease in tissue viability is < 30%. The PM resulted in high sensitivity (91.7%) and specificity (100.0%) for 42 test materials (24 phototoxic/18 non-phototoxic). Results of *iPHO* assay were compared to in-house 3T3-NRU-PT assay.

The current protocol extends phototoxicity testing using EpiDerm for risk assessment to systemically administered chemicals and medications and will provide the pharmaceutical industry with an in vitro screening method to assess the phototoxic risk of new API.
Assessing Eye Irritation Potential of Cosmetic Products Using the STE Test.

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Various in vitro assays that have been developed as an alternative for the Draize test are currently used to evaluate the eye irritation potential of cosmetic products and ingredients. A Short Time Exposure (STE) test, which we developed as a potential alternative test for eye irritation using rabbit cornea cells (SIRC), is planned for peer review and expected to be accepted as an OECD test guideline for classifying materials as Non-irritant/Irritant (NI/I). The test is expected to be able to evaluate cosmetic products as well as chemicals and cosmetic ingredients. Firstly, we evaluated the accuracy of the STE test against GHS classification for 20 cosmetic products (14 rinse-off and 6 leave-on products). Secondly, the results of the STE test (including various rinse-off and leave-on products) were compared with the results of 43 cosmetic products tested in BCOP (bovine cornea opacity and permeability) test, 20 cosmetic products tested in HET-CAM, and 40 cosmetic products tested in CAMVA (two hen egg chorioallantoic membrane tests) in order to confirm the homology of in vitro assays used for evaluating eye irritation of the cosmetic products. In the first step, the accuracy for STE classifying a product as NI/I, compared to GHS, was 90% (18/20). Furthermore, STE test is able to provide a rank order (minimal irritant, moderate irritant and severe irritant) for materials based on the cell viability. The accuracy which compared GHS classification to a rank order of STE test was 75% (15/20). In the second step, comparison between STE test and three in vitro assays demonstrated that the accuracy of STE classifying a product as NI/I compared to BCOP, HET-CAM, and CAMVA was 79% (34/43), 80% (16/20), and 90% (36/40), respectively. Our findings show that STE test can be an alternative for the Draize test as well as for the three in vitro assays used to evaluate a cosmetic product.

Choosing the Appropriate Solvent for Solid Materials Tested in the Bovine Cornea Opacity and Permeability (BCOP) In Vitro Assay.

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In compliance with OECD Test Guideline 437 for eye irritation (BCOP) assay, non-surfactant solid materials are typically tested as 20% dilutions prepared in 0.9% sodium chloride solution, distilled water, or other solvent that has been demonstrated to have no adverse effects on the test system. However, the limited solubility of some chemicals adds technical challenges in finding a vehicle that would ensure the material’s availability to the excised corneas and that itself would not affect the test system. In this study, we evaluated five solvents frequently used in the BCOP assay: distilled water, mineral oil, corn oil, polyethylene glycol (PEG-400), and methocel solution (0.5%). Based on the available classification systems, our preliminary data showed that water, methocel, mineral oil and corn oil were predicted as non-irritants, while PEG-400 was predicted as a mild irritant. To demonstrate the influence of the type of solvent on the outcome/prediction of the BCOP assay for solid materials, we tested a 20% suspension of benzoic acid (BA) prepared in these solvents. BA has a non-polar benzoic ring that would preferably dissolve in non-polar solvents and a polar acidic group with affinity for polar solvents, thus making it a good model for testing its effect on corneas when dissolved in various solvents. Previous animal tests reported moderate to severe eye irritation induced by BA. Our results demonstrated that when mixed in water, mineral oil, corn oil, or methocel, BA was predicted to be a corrosive/severe irritant, while it was predicted to be a moderate irritant when mixed in PEG-400. These results support the need for further investigation of the solvent’s influence in the BCOP assay to allow the correct prediction of the irritation potential of solid materials.
The bovine corneal opacity and permeability (BCOP) test has been adopted by OECD for the identification of ocular corrosive and severe irritants (GHS category 1) for single component substances and multi-component formulations. Further, human reconstructed tissue models have been suggested for incorporation into a tiered test strategy to ultimately replace the Draize rabbit eye irritation test (OECD TG 405) and the value of the EpIcular Eye Irritation Test (EIT) for the prediction of ocular non irritants (GHS no category) has been shown previously. The purpose of this study was to evaluate whether the BCOP including corneal histology and the EIT could be used to predict eye irritancy of agrochemical formulations according to different classification schemes including UN GHS, EPA and ANVISA systems. We have assessed opacity, permeability and histology in the BCOP assay and relative tissue viability in the EIT for the 50 agrochemical formulations with available in vivo eye irritation data. Using the OECD TG guideline evaluation scheme for opacity and permeability in the BCOP did not prove predictive with respect to severe eye irritation potential for the 50 agrochemical formulations assessed here, while corneal histology grades and the EpIcular tissue viabilities where useful predictors of eye irritancy potencies. Further we describe here statistical evaluation based on the experimental in vitro data to predict eye irritancy for the different classification schemes.

A Preliminary Investigation in the Use of Porcine Corneas As a Substitute for Bovine Corneas in the BCOP Assay.

D. Wolffinger and D. R. Cerven. MB Research Laboratories, Spinnerstown, PA. Sponsor: G. DeGeorge

The bovine Cornea Opacity and Permeability Assay (BCOP) has been used since 1992 to provide estimates of ocular irritation. It has been accepted as an alternative to the use of rabbit in acute ocular safety testing. However in some areas of the world, access to bovine eyes is limited, and the Draize eye-scoring test, performed in rabbits, is still being utilized. In response to this, we have been investigating the replacement of excised bovine corneas with porcine corneas for evaluation of acute ocular irritation (PCOP). This preliminary investigation involved the evaluation of 100% ethanol, 0.1% Benzalkonium Chloride (BAK), and four cosmetic products, all of which had previously been tested in the BCOP as well as the Chorioallantoic Membrane Vascular Assay (CAMVA). Initially the results from the ethanol and BAK testing in the porcine corneas showed no correlation to the results from bovine corneas. These suggested the need for certain modifications to the trimming and mounting procedures for the porcine corneas, which are smaller than the bovine corneas. Thus, the trimming process to excise the cornea was modified, and the same chambers supplied with the opacimeter for use with bovine eyes was also used for the porcine eyes. After modifying the pre-testing procedures, we proceeded to test four cosmetic compounds successfully in the BCOP and PCOP. These investigations indicate that the In Vitro Scores obtained with porcine eyes are similar to those in bovine eyes. Additional evaluation with a wider range of chemical entities is underway.

In Vitro Ocular Irritation Testing Strategy for Prototype Cleaning Products.

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The Bovine Corneal Opacity and Permeability (BCOP) assay can be used for predicting mild, moderate, and severe ocular irritation through quantitative assessment of the changes in opacity and permeability of bovine corneas. In addition, histological evaluation of the corneas may be performed to assess the depth of damage. The BCOP assay with histology was used to determine the ocular irritation potential of prototype cleaning products with antimicrobial claims according to the guideline provided by the EPA-Office of Pesticide Programs (OPP). Several prototype cleaners of similar formulation were evaluated along with a reference material. The results of the BCOP assay showed noticeable differences among the products. The in vitro score, determined by changes in opacity and permeability of the corneas, ranged from ~15 to 80. These scores indicate mild, moderate, and severe irritation according to the guideline provided by the EPA-OPP. In addition, the histological evaluation of the corneas showed differences in the depth of damage to the stroma between moderate and severe category products, confirming the classification of the products by the in vitro score. These results demonstrate the utility of the BCOP assay with histology as a stand-alone assay for eye irritation evaluation in the EPA-OPP program. Furthermore, this testing strategy distinguished the ocular irritation potential among similar prototypes demonstrating its effectiveness in the product development process.


L. d’Argenbeau-Thornton, Y. Kaluzhny, H. Kandarava, P. Hayden and M. Klaunzer. MatTek Corporation, Ashland, MA.

The FDA and other regulatory agencies require ocular irritation testing of ophthalmologic pharmaceuticals and consumer products. Human reconstructed tissue models have been suggested for incorporation into a tiered testing strategy to replace the Draize eye irritation test (OECD TG 405) and for use as highly reproducible and cost effective in vitro alternatives. A newly developed Ocular-Corneal Full Thickness (OCFT) tissue model includes epithelial, stromal, and endothelial layers. This model has been characterized in terms of tissue morphology, expression of specific markers for human corneal epithelium (Cytokeratins 12, 14, and 19, stratfin, tight junction proteins ZO-1, occludin, and claudin), and Na/Cl leakage and transepithelial electrical resistance (TEER) in response to chemical insults. Histological evaluation of the tissues was used to provide information on the type and depth of the chemical insult. Also, the OCFT model was used to assess tissue recovery following exposure to increasing concentrations of surfactants and TA spanning the range of irritancy. Significant recovery was observed in 24 and 48 hours for non-irritants (NI) chemicals. The OCFT tissue was also evaluated in a response to a battery of test articles (TA, n=65) spanning the range from ocular corrosives and severe irritants (I) to non-irritants (NI). In vivo irritancy levels were obtained from a published database and in vitro effects were determined using the MTT tissue viability assay and TEER. Using a tissue viability cutoff of 50%, the OCFT was able to detect “I” with 100.0/75.0/and 87.7% specificity/sensitivity and accuracy. The OCFT model will address the needs of ophthalmic and other formulators who need to screen their products for irritancy and efficacy. Similar to other in vitro assays, high levels of reproducibility at a low cost are anticipated. The OCFT model will facilitate the development and testing of ophthalmologic pharmaceuticals and allow for basic studies related to corneal physiology, wound repair, and pathologies.

Validation of In Vitro Eye and Skin Irritation Assays for Structurally Complex Pharmaceutical Chemicals.

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Identification of local ocular and dermal irritation effects is important for occupational health and hazard communication. OECD has published Test Guidelines (TG) 439 for In Vitro Skin Irritation (EpiDerm™ System, MatTek) and TG 437 for the Bovine Corneal Opacity and Permeability Test Method (BCOP) to replace or supplement in vivo tests, but their applicability to complex molecules found in the pharmaceutical industry has not been fully investigated. A validation study was conducted to compare in vitro assays (OECD TG 405) and skin irritation (OECD TG 404) to in vivo tests for data for Bristol-Myers Squibb (BMS) proprietary compounds. The validation set consisted of 15 compounds, which were selected to represent the BMS chemical space within six chemical classes: α/unsaturated carbonyls, α-halo ketones, aminoaacohols, boronic acids/esters, halogenated heterocycles, and nitroaromatics. The assay results were interpreted using OECD and GHS classification schemes and in vitro and in vivo results were compared using basic statistics. The BCOP assay accurately predicted 12 of 15 compounds when non- and mildly irritating classifications were grouped together. Both boronic compounds were under-predicted by BCOP, suggesting the assay may not be appropriate for this chemical class. The in vitro skin assay demonstrated high specificity by correctly identifying 12 of 13 negative in vitro skin irritants, but, the sensitivity could not be interpreted from the limited number of positive skin irritants in this study. Additional data are needed to assess the predictivity of positive skin irritants with greater confidence. Overall, the replacement of in vivo studies with these alternative in vitro irritation methods reduces and refines the tiered testing strategy for ocular and dermal hazard identification of pharmaceutical chemicals.

In Vitro Ocular Irritation Testing Methods.

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The results of the BCOP assay showed noticeable differences among the products. The BCOP assay with histology was used to determine the ocular irritation potential of the changes in opacity and permeability of bovine corneas. In addition, histological evaluation of the corneas showed differences in the depth of damage to the stroma between moderate and severe category products, confirming the classification of the products by the in vitro score. These results demonstrate the utility of the BCOP assay with histology as a stand-alone assay for eye irritation evaluation in the EPA-OPP program. Furthermore, this testing strategy distinguished the ocular irritation potential among similar prototypes demonstrating its effectiveness in the product development process.


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A newly developed Ocular-Corneal Full Thickness (OCFT) tissue model includes epithelial, stromal, and endothelial layers. This model has been characterized in terms of tissue morphology, expression of specific markers for human corneal epithelium (Cytokeratins 12, 14, and 19, stratfin, tight junction proteins ZO-1, occludin, and claudin), and Na/Cl leakage and transepithelial electrical resistance (TEER) in response to chemical insults. Histological evaluation of the tissues was used to provide information on the type and depth of the chemical insult. Also, the OCFT model was used to assess tissue recovery following exposure to increasing concentrations of surfactants and TA spanning the range of irritancy. Significant recovery was observed in 24 and 48 hours for non-irritants (NI) chemicals. The OCFT tissue was also evaluated in a response to a battery of test articles (TA, n=65) spanning the range from ocular corrosives and severe irritants (I) to non-irritants (NI). In vivo irritancy levels were obtained from a published database and in vitro effects were determined using the MTT tissue viability assay and TEER. Using a tissue viability cutoff of 50%, the OCFT was able to detect “I” with 100.0/75.0/and 87.7% specificity/sensitivity and accuracy. The OCFT model will address the needs of ophthalmic and other formulators who need to screen their products for irritancy and efficacy. Similar to other in vitro assays, high levels of reproducibility at a low cost are anticipated. The OCFT model will facilitate the development and testing of ophthalmologic pharmaceuticals and allow for basic studies related to corneal physiology, wound repair, and pathologies.
and H. are often more sensitive to respiratory viral infections later in life. Similarly, adult oxygen (hyperoxia) at birth are at increased risk for impaired lung development and adult diseases. For example, preterm infants prematurely exposed to excess levels of virus infection, which may provide insight into how early-life exposure to high levels of oxygen leads to childhood and adult diseases later in life.

Late lung development is a critical window of susceptibility as perturbations impair alveolarization and vascular development. Currently, genetic control of differential gene expression during lung development and injury is incompletely understood. We hypothesized that integration of genetics and genomics in expression quantitative trait loci (eQTL) analysis will identify genetic factors contributing to variability in the neonatal lung and characterize genetic susceptibility. Full term neonates from 30 inbred strains of mice were maintained in room air or exposed to 95% oxygen for 72 hours beginning on postnatal day 1 (P1). Lungs were collected from 3 neonates/group for each strain on P4. RNA was isolated for transcriptomic analysis using Illumina WG6v2.0 BeadChips. Transcript intensities were analyzed for differential expression, and associations between genetic polymorphisms and transcript intensity (eQTLs) were identified by haplotype association mapping with FastMap. 4513 differentially expressed transcripts were identified on P4, and cis-eQTL analysis, or associations between a transcript and polymorphism in the same gene, yielded 29 genes. Biologically interesting candidate genes included Adam17, Pdx1, Tnfalpha2, and Eif2ak2. After hyperoxia exposure, 5386 transcripts were differentially expressed. 1230 were specifically hyperoxia-induced and related to developmental and inflammatory pathways. 23 cis-eQTLs were identified and biologically interesting candidates included Vep, Cas9, and Trim30. These analyses identified candidates with transcription levels in the developing lung that is, in part, due to genetic background. Furthermore, network and pathway analysis of these candidate genes pointed to a novel role for the integrated stress response and proteostasis in neonatal lung development and injury.

Environmental exposures combined with genetics can have a persistent influence on common chronic health conditions of children and adults. A growing body of research at the National Institute of Environmental Health Sciences (NIEHS) is focusing on environmental effects during early-life development and disease occurrence over the lifespan—a branch of scientific study referred to as Developmental Origins of Health and Disease or the “DOHaD paradigm.” This paper describes the evolution of DOHaD research at NIEHS and how extramural events, including Requests for Applications (RFA) in the areas of Fetal Basis of Adult Disease (FeBAD), children’s health, and epigenetics, have stimulated the field of developmental origins of health and disease. A growing body of research at the National Institute of Environmental Health Sciences (NIEHS) is focusing on environmental effects during early-life development and disease occurrence over the lifespan—a branch of scientific study referred to as Developmental Origins of Health and Disease or the “DOHaD paradigm.” This paper describes the evolution of DOHaD research at NIEHS and how extramural events, including Requests for Applications (RFA) in the areas of Fetal Basis of Adult Disease (FeBAD), children’s health, and epigenetics, have stimulated the field of developmental origins of health and disease. A growing body of research at the National Institute of Environmental Health Sciences (NIEHS) is focusing on environmental effects during early-life development and disease occurrence over the lifespan—a branch of scientific study referred to as Developmental Origins of Health and Disease or the “DOHaD paradigm.” This paper describes the evolution of DOHaD research at NIEHS and how extramural events, including Requests for Applications (RFA) in the areas of Fetal Basis of Adult Disease (FeBAD), children’s health, and epigenetics, have stimulated the field of developmental origins of health and disease. A growing body of research at the National Institute of Environmental Health Sciences (NIEHS) is focusing on environmental effects during early-life development and disease occurrence over the lifespan—a branch of scientific study referred to as Developmental Origins of Health and Disease or the “DOHaD paradigm.” This paper describes the evolution of DOHaD research at NIEHS and how extramural events, including Requests for Applications (RFA) in the areas of Fetal Basis of Adult Disease (FeBAD), children’s health, and epigenetics, have stimulated the field of developmental origins of health and disease. A growing body of research at the National Institute of Environmental Health Sciences (NIEHS) is focusing on environmental effects during early-life development and disease occurrence over the lifespan—a branch of scientific study referred to as Developmental Origins of Health and Disease or the “DOHaD paradigm.” This paper describes the evolution of DOHaD research at NIEHS and how extramural events, including Requests for Applications (RFA) in the areas of Fetal Basis of Adult Disease (FeBAD), children’s health, and epigenetics, have stimulated the field of developmental origins of health and disease. A growing body of research at the National Institute of Environmental Health Sciences (NIEHS) is focusing on environmental effects during early-life development and disease occurrence over the lifespan—a branch of scientific study referred to as Developmental Origins of Health and Disease or the “DOHaD paradigm.” This paper describes the evolution of DOHaD research at NIEHS and how extramural events, including Requests for Applications (RFA) in the areas of Fetal Basis of Adult Disease (FeBAD), children’s health, and epigenetics, have stimulated the field of environmental health science as it relates to DOHaD. In order to build and analyze a portfolio of active NIEHS-funded grants in the area of DOHaD, the electronic Scientific Portfolio Assistant (eSPA) software application was used. DOHaD portfolios for the years 1991, 2001, and 2011 were analyzed for exposures, disease/organ endpoints, windows of exposure, study design, and impact on the field based on publication data. Priority exposures in the 1991 and 2001 portfolios comprise of metals (e.g., lead and mercury), PCBs, and air pollutants; but by 2011, the portfolio has evolved to include or expand the variety of endocrine disruptors (e.g., BPA, phthalates), pesticides/Persistent Organic Pollutants (POPs) (e.g.,.
organophosphates, organochlorines, PBDE, PAH, and metals (e.g. arsenic, man- ganese). Several disease endpoints related to brain/CNS are apparent across all three portfolios, whereas reproduction and cancer increase steadily over the same time period, and new endpoints like obesity are introduced by 2011. The analysis of NIEHS-funded DOHaD research provides insight into the institute’s impact on the field, and will help determine how to improve the quality and health impact of future, funded DOHaD projects.

987 Effects of Gutkha Use during Pregnancy on Hepatic Parameters Later in Life.

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Gutkha, a popular smokeless tobacco (ST) herbal concoction made in India and sold throughout India and the U.S., is a combination of tobacco, lime, betel and areca nut, spices, and catechu. Recent evidence demonstrates that consuming gutkha during pregnancy may be linked to adverse obstetric consequences. However, whether gutkha exposure during pregnancy increases offspring risk of chronic disease later on in adulthood, as with combustible tobacco products, remains unknown. For this study, pregnant B6C3F1 mice were exposed daily to 25 mg of a water-soluble gutkha solution beginning on gestational day (GD) 2-4 until parturition. The main objectives of this study were to evaluate prenatal exposure to gutkha altered offspring susceptibility for hepatic disease later in life. After male, female, mice were exposed to a gutkha solution via the oral mu cosa. Serum cotinine levels in pregnant mice were measured weekly and ranged from 24-44 ng/mL. Cotinine levels, also measured in the amniotic fluid and fetal liver on -GD 15, averaged 21-22 ng/mL. Beginning at 12-weeks of age, a sub-group of male and female offspring were fed a high fat (HF) diet for 14 days. Male offspring prenatally exposed to gutkha and fed subsequently a HF diet had increased hepatic fat mass, fibrosis, and inflammation compared to the other groups. Additionally, female offspring exposed to gutkha alone had significantly higher serum levels of the hepatic injury markers, alanine transaminase and aspartate aminotransferase. In conclusion, findings from these studies demonstrate that gutkha usage during pregnancy may have persistent later life effects that increase the risk of adult liver disease. As gutkha use by pregnant women in the South Asian communities is increasing exponentially, such a menacing public health behavior requires im- mediate attention. Supported by Memorial Sloan Kettering Cancer Institute.

988 Transcriptomic Profiling of Embryos and Adult Female Brain Tissue Links a Developmental Origin of Atrazine-Induced Reproductive Alterations in Zebrafish.

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Atrazine, an herbicide commonly applied to agricultural areas throughout the Midwest and a common contaminant of potable water supplies, is implicated as an endocrine disruptor and potential carcinogen. The specific adverse health effects asso ciated with atrazine exposure and the underlying molecular mechanisms of these effects are not well defined. In an effort to delineate the mechanisms of atrazine toxicity, we exposed zebrafish embryos to environmentally relevant concentrations of atrazine shortly after fertilization through 72 hours post fertilization (hpf). Transcriptomic profiles immediately following the embryonic atrazine exposure were obtained and subsequent analyses identified an enrichment of genes with altered expression patterns that are involved in neuroendocrine development and function, cell cycle regulation and carcinogenesis. From these exposures a subset of individuals was permitted to mature under normal conditions to evaluate later life effects of the developmental exposure and in unexposed subsequent generations. A significant difference in the number of pairs that successfully bred in one of the atrazine treatment groups coupled with distinct altered reproductive histological and morphological alterations in female adult zebrafish was observed indicating a later life effect of the developmental atrazine exposure. Furthermore, gene ontology analysis from microarrays performed on adult female zebrafish brains showed enrichment for genes involved in neuroendocrine system function and disease. The reproductive alterations observed in the adult females along with decreased mating success and altered transcriptomic profiles provide support to the endocrine disrup- ting effects of this herbicide and warrant further mechanistic investigation. Current efforts are aimed at assessing transgenerational effects of this developmental atrazine exposure in subsequent unexposed generations.

989 Bisphenol A and Diethylstilbestrol Treated Mice Respond Differently to a Very High Fat Diet.

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Rationale: Bisphenol A (BPA) is an estrogenizing endocrine disruptor, and diethylstilbestrol (DES) is a non-steroidal estrogen. Some data suggest that BPA is an obe- sogen. We hypothesized that BPA and/or DES would increase the metabolic impact of a very high fat diet and that this obesity would reduce heart structure/function. Methods: C57bl/6J mice were exposed to Vehicle, oral BPA (5.0 μg/kg/da) from gestation day 11 to weaning and then (0.5 μg/kg/da) to 4 months, or diethylstilble- strol (DES, 1μg/kg/da) from GD 11-14. Mice were fed control (CD, 10% calories from fat) or very high fat (HFD, 60% calories from fat) diets from weaning. Body weight (BW) and body length (BL) were measured monthly. Body mass index (BMI) and body surface area (BSA) were calculated. Glucose tolerance was tested at 15 weeks. Echocardiography, heart weights and serum for cholesterol, triglycerides and leptin were collected at 16 weeks. Results: All HFD mice had increased BW, BL, BMI and BSA. HFD VEH and BPA males had similar increases in total fat and total fat indexed to BW or BSA, but HFD BPA males were smaller in BW, BMI and BSA. HFD VEH and BPA females had similar total fat and total fat indexed to BW or BSA, BMI and BSA. In contrast, HFD DES treated males and females had the least fat gain, BW, BMI and BSA. Cholesterol and triglycerides were increased with HFD in all males and leptin was increased in BPA and DES males. Cholesterol, triglycerides and leptin were in- creased with HFD identically in VEH, BPA and DES females. Glucose tolerance was reduced with HFD equally in all mice. Fractional shortening was unaffected, but HFD induced eccentric cardiac hypertrophy in VEH and BPA with DES mice showing the greatest increase. Conclusions: Neither BPA nor DES worsened the response to a very high F. VEH and BPA mice responded similarly to a very HFD. DES mice had a blunted meta- bolic response to a very HFD. Cardiac function was unaffected, but cardiac struc- ture was affected by HFD.

990 Analysis of Transcriptomic Profiles and Functional Clustering of Global Cardiovascular Gene Expression in Response to In Utero B(a)P Exposure.

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Interest in the correlation between environmental toxicants and cardiovascular dis- eases has increased considerably over the past few decades. However, little is known of B(a)P’s effect on the developing heart or the specific biological pathways altered by B(a)P exposure. Functional comparative genomic analysis of the cellular and development- al effects of different polycyclic aromatic hydrocarbon (PAH) compounds is con- sidered a helpful approach to distinguish the complex and specific effects of B(a)P exposure in utero. Using timed-pregnant Long Evans Hooded rat offspring, we characterize the genetic modulation in cardiovascular related genes for three concentra- tions of B(a)P (0, 600, and 1,200 μg/kg/BW) at postnatal days 0 and 53 (P- 0 and P-53). In utero exposure to B(a)P at both 600 and 1,200 μg/kg/BW significa- ntly modulated the expression of 89 out of 45,000 genes in offspring. The focused microarray approach identified important subgenomic differences in the pattern of cardiovascular disease cell-related gene expression in response to in utero B(a)P exposure. These molecular targets and deduced networks may be employed as a guide for classifying, monitoring and manipulating the molecular and patho- logical specificities of different PAHs in key cardiovascular related cell systems and for potential pharmacological application.

991 Maternal DDT Exposure Increases Risk of Metabolic Syndrome in Adult Offspring.

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DDT is associated with altered carbohydrate and lipid homeostasis in animals that have been exposed to relatively high doses, and with increased risk of type 2 dia- betes and hypertension in human studies. We hypothesized that maternal exposure
to DDT would increase the risk of metabolic syndrome in the adult offspring of mice. We administered DDT to C57BL6J mice from gestational day 11.5 to postnatal day 5, when litters were culled to six pups. We performed comprehensive metabolic phenotyping in offspring. Maternal DDT exposure increased diastolic- and systolic-blood pressure in adult offspring. Maternal DDT exposure also caused a mild obese phenotype in female offspring that was due to impaired thermoregulation (increased cold intolerance) and decreased energy expenditure while food intake was not different. When these female offspring were fed a diet high in fat, they developed elevated fasting insulin and lipids. Thus, the results from the current study suggest that maternal DDT exposure leads to metabolic syndrome in adult offspring.

PS 992 Mechanisms Underlying Neurodevelopmental Defects in Weanling and Adult Rats Associated with Prenatal Exposure to Methylmercury.

D. De Groot1, C. De Esch1, M. de Groot1, C. Kuper1, R. Steur1, A. Wolterbeek1, E. De Vries2, N. Cappaert3, M. Radonjic1 and R. TBTOdev on brain SG was most pronounced on cerebrum at PD22, which was no PD22, whereas at PD62 both weight and volume appeared reduced. Effects of juvenile exposure appeared mild, and seemed to affect both cerebrum and cerebellum/brainstem (PD62). The results are discussed in light of TBTO-induced delayed development, persistent effects at adult age and a speculative role of TBTO as obesogen.

PS 994 The Effects of Endocrine Disruption on the Maturation of the Developing Human Fetal Prostate.

C. Safarini1, E. V. McDonnell1, A. Amin1, S. J. Hall1 and K. Boekelheide1.

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The etiology of prostate cancer is unknown, although it has been suggested that early life exposures to various toxicants, such as estrogenic chemicals, may play an important role. Previous studies in rat models have demonstrated that early life exposures to estrogens is responsible for causing epithelial and stromal hyperplasia, inflammation, and prostatic intraepithelial neoplasia (PIN) lesions. We have developed a xenograft rodent model to characterize the growth and differentiation of human fetal prostate implants (gestational age 12-24 weeks). This model encompasses the effects of both early and later-life exposures to either corn-oil (control) or 250 μg/kg/bone weight of 17β-estradiol 3-benzoate post-transplant. To ensure proper growth of implants, the renal subcapsular space was chosen as the site of implantation to allow for appropriate growth and vascularization of prostate tissue. This model was characterized based on the expression of key immunohistochemical markers responsible for epithelial and stromal maturation including p63, vimentin, α-smooth muscle actin, caldesmon, ki-67, prostate specific antigen, estrogen receptor-α, and androgen receptor. As expected, the human fetal implants grew and matured as demonstrated by the histopathology seen in 7, 14, 30, 90, 200 and 400 day xenografts. Interestingly, the human prostate xenografts exhibited marked differences in response to estrogen exposure compared to their endogenous rodent prostate counterparts. The endogenous rodent prostates exhibited a mild obese phenotype at PD62 as indicated by p63 staining. This study of human fetal prostate tissue will allow for future mechanistic studies investigating the origins of prostate disease.

PS 995 Characterizing Later-Life Spatial Discrimination Deficit Phenotypes following In Utero Exposure to Benzoprene.

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To characterize observed behavioral learning deficit phenotypes in offspring adult rodents arising from in utero exposure to benz(a)pyrene (B(a)P), timed-pregnant Long Evans Hooded rats or Cytochrome p450-2e1 (Cyp2e1) mice were exposed to either B(a)P (0, 150, 300, 600 and 1200 ng/kg BW or aerosol 100 μg/m3 on embryonic days 14 to 17. In the pre-weaning period, B(a)P metabolites were examined in cerebral cortex. No detectable levels of metabolites were found in the control offspring during the pre-weaning period but dose-related increases in B(a)P metabolites were seen in cerebral cortex and revealed a time-dependence for elimination during this period. Reversal learning was evaluated using a spatial discrimination-reversal procedure in adult offspring. Offspring that were exposed in utero to 600 and 1200 μg/kg BW or 100 μg/m3 required a significantly higher number of sessions to complete the first reversal than control offspring. However, with original discrimination and subsequent reversals, exposed offspring behavioral profiles resembled their respective controls. This effect was seen in errors of commission, indicative of perseveration or resistance to extinction. Offspring that were exposed in utero also had shorter choice latencies than controls during the first session of a reversal. Activity-related-cytoskeletal associated protein (Arc), an experience-dependent cortical protein marker known to be up-regulated in response to acquisition of novel behaviors, was also upregulated in B(a)P exposed animals that were trained in the behavioral paradigm. Collectively, the findings support the hypothesis that in utero exposure to B(a)P during the critical window for peak neurogenesis produces later-life learning deficit phenotypes in offspring.
The continued development of engineered nanomaterials (ENM) has given rise to concerns over the potential for human health effects. While the lung is the primary site of exposure, the cardiovascular system is a principal site of impact. One of the most complex and acutely demanding circulations is the enhanced maternal system to support fetal development. The “Barker Hypothesis” proposes that metabolic impairments during gestation predispose future senescence. Herein we initiated testing at the microvascular level. Pregnant (gestation day 10) Sprague-Dawley rats were exposed to nano-titanium dioxide aerosols (count mode aerodynamic diameter of 137 nm, 10 mg/m³, 4 hrs/day) for 10 days to evaluate the maternal and fetal consequences of maternal exposure. The calculated daily maternal deposition was 20 μg. These exposures lead to fetal irregularities and maternal microvascular dysfunction. Isolated maternal uterine arteriolar (< 150 μm) reactivity was significantly impaired overall, consistent with a metabolically impaired profile. This dysfunction presented as blunted endothelium-dependent reactivity (acetylcholine, ACh, 10-9-10-4 M), reduced adrenergic responsiveness (phenylephrine, 10-9-10-4 M), and impaired myogenic responsiveness (transmural pressure, 15-120 mm Hg). With respect to the fetal group, maternal exposures lead to a significant decrease in total fetal weight. Fetal tail arteries (< 150 μm) were isolated to assess microvascular alterations after maternal exposure. Interestingly, the vessels from the exposed group also demonstrate significant impairment to increasing concentrations of ACh (10-9-10-4 M), spermine NONOate (10-9-10-4 M), and increases in transmural pressure. Collectively, impaired uterine microvascular reactivity after ENM exposure during pregnancy can reduce nutrient exchange, drastically impacting fetal weight and number. This prenatal exposure may also lead to cardiovascular consequences for the developing fetus. NIH-R01-ES015022 and RC1-ES018274 (TRN) NSF-1003907 (VCN)

The AhR Contributions to Cardiovascular Development, Developmental Toxicity, and Adult Disease.

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Developmental disruption has been increasingly recognized as a key contributor to adult disease. Exposures to ubiquitous environmental persistent organic pollutants (POPs), which includes the pervasive organic pollutant 2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD), can result in abnormalities of the cardiovascular system at the structural and functional levels. TCDD is a prototypical aryl-hydrocarbon receptor (AhR) ligand but the precise pathogenesis and phenotype of TCDD developmental toxicity remains poorly characterized. Our work aims at understanding how developmental perturbations (TCDD-exposure) or absence (knockout) of the AhR pathway affects the cardiovascular system in the embryo and adult. To characterize the toxicity phenotype, C57BL/6J pregnant dams were oral-gavaged with different doses of TCDD (0.1, 0.5, 1, 2.5, 5, and 50 μg/Kg) or vehicle control at key murine cardiogenesis time points (E7.5, 9.5, and 11.5). Embryos were harvested at E13.5, 15.5, and 18.5 or carried to term and adulthood. Additionally, C57BL/6J Ahr-/- mice were included in postnatal functional studies. At all studied doses, embryos could be carried to term; however, pups died within 24 hours post-partum at the three highest doses. Embryo cardiac toxicity was characterized by dynamic changes in morphometric parameters (right and left ventricular free wall thickness, cellular density, nuclear size, and extracellular matrix) as well as the changes in the expression of the AhR and cellular proliferating proteins. The postnatal functional studies highlighted LV cavity dilatation in the Ahr-/- mice at PDN60 and 90. These results corroborate the heart as a target of developmental disruption by in-uterus exposure to a POP. Moreover, since all biological effects of TCDD exposure are believed to be mediated through AhR signaling, these results underscore the critical role for the AhR pathway in normal cardiovascular development. Supported by NIH Grant R01ES050673.
Embryonic vascular development can be disrupted by diverse compounds (thalidomide, estrogen, nicotine, dioxin, retinoids, cigarette smoke, metals). Collaborative studies are underway to test these predictions across an array of assays (whole embryo culture, aortic explants, endothelial tubulogenesis, transgenic zebrafish, etc.). Here, we describe initial results with a complex human-cell based in vitro angiogenesis assay, on the first 7 of 20 ToxCast Phase I chemicals selected by a range of predicted activity as vascular disruptors (pVDCs). Degree of endothelial tubule formation in vitro was scored by AC50 values relative to cytotoxicity. Both chemicals predicted as non-pVDCs, imazamox and pyremethrin, showed no inhibitory activity. The predicted inhibitory potential of 5 pVDCs was confirmed for 4 out of 5 chemicals. Pyridaben, predicted as a strong pVDC, inhibited tubule formation in a concentration-dependent manner in the FICAM assay, first evident at 0.001 μM and complete at 0.01 μM. Other predicted pVDCs with different ToxCast activity profiles (BPA, oxytetracycline, fluazinam) inhibited endothelial tubulogenesis at concentrations ranging from 5- to 500 μM. PFOS was the only active pVDC tested; however, the ToxCast prediction was based on inhibition of Ang1/Tie2 signaling that controls later stages of vessel stabilization— a subtle outcome in the ABM simulation. Overall, concordance between the tubulogenesis assay and computer ABM simulation demonstrates the utility of these in vitro and in silico models for predictive modeling and mechanistic understanding of vascular disruption in higher-order biological tissues. [This work does not reflect EPA policy].

1003 Placental Transfer of 125I-iodinated Humanized Immunoglobulin G2Aα in the Sprague Dawley Rat.

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Biopharmaceuticals are a growing segment of the health care therapeutic arsenal and indications for these agents are rapidly expanding into patient populations that include women of child bearing potential. The fetus is typically not a target of these drug therapies, and thus effects on the developing conceptus are generally undesirable. While small molecules generally cross the placenta via passive diffusion throughout gestation, transfer of large molecules is limited and may be species dependent. Antibody-like biopharmaceuticals (Abs), that contain an Fc region, are unique and cross the placenta using active transport pathways normally reserved for maternal immunoglobulin G (IgG). Embryo/fetal, placental and maternal tissue concentrations of an Ab in Sprague Dawley rats were evaluated by 2 different biodistribution techniques following maternal injection of 125I-iodinated IgG2Aα on different gestation days. Blood and tissue samples were collected for gamma counting or whole maternal carcasses were processed for quantitative whole body autoradiography. The results indicate the presence of maternally injected IgG in the conceptus as early as Gestation Day (GD) 11 and a >1000 fold increase in that amount by GD 21; and maternalAb concentrations generally remaining unchanged during gestation (within the same order of magnitude). In addition, fetal/maternal tissue concentration ratios remain stable during organogenesis with only a slight increase at the end of gestation. These data indicate that Abs with a target present in the developing conceptus have the potential to elicit an unintended biological response depending on the Ab affinity and potency. These data also demonstrate that maternally-administered Abs may be present during organogenesis and have the potential for adverse developmental outcomes in the rat based on direct embryo/fetal exposure.

1004 Evidence That Perfluoroaoctic Acid (PFOA) Does Not Activate Estrogen Receptor (ER) Activities In Vivo or In Vitro.

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We evaluated the potential of perfluorooctanoate to produce estrogenic activity in both an in vivo utero-sac assay and an in vitro estrogen receptor (ER) activation assay. In the in vivo study, pre-puberal female CD-1 mice born to dams fed an estrogen-free diet from PND 14 through weaning on postnatal day (PND 18) were given daily oral doses of 0.005, 0.01, 0.02, 0.05, 0.1 or 1 mg/kg PFOA on PND 18-20. In addition, a positive control received daily 17β-estradiol (E2, 0.5 mg/kg). On PND 21, the uterus and vagina were weighed and prepared for histology. Transcripts concentrations for the ER responsive genes P21, progesterone receptor (PR), and trefoil factor (TFF) were detected and quantified in the uterine tissue using real-time quantitative RT-PCR. Serum PFOA concentrations were determined by LC-MS/MS. PFOA caused no changes in body weights, uterine weights, transcripts for ER target genes, or in squamous hyperplasia and cornification of the vaginal epithelium. ER administration produced increased relative uterine weight, increases in transcript concentrations for ER-responsive genes, and squamous hyperplasia and cornification of the vaginal epithelium. Serum PFOA ranged from 5 to 1200 ng/mL. In the in vitro study, PFOA at 0.0005, 0.001, 0.005, 0.01, 0.05, 0.1, 5 or 10 μg/mL in media failed to affect estrogen response element activity in a human ovarian carcinoma cell line (BG1-Luc/E2) that carried an estrogen response element (ERE)-luciferase reporter construct; whereas, E2 exhibited a marked increase. Collectively, these data do not support an estrogenic effect of PFOA either in vivo or in vitro.
1005 Effects of Di(2-Ethylhexyl)-Phthalate Exposure during the Periparturient Period: An Analysis of the Evidence.
Di(2-ethylhexyl)phthalate (DEHP) is a widely-used plasticizer and ubiquitous environmental contaminant. DEHP exposure during late gestation in rodents has been shown to cause a spectrum of adverse effects related to altered androgen function, collectively termed the ‘phthalate syndrome’. Thus, many studies on DEHP have focused on exposures during this critical period of organ development and outcome measures in male offspring. A more limited database exists for other endpoints or following exposures that do not encompass fetal development. One such exposure paradigm involves DEHP treatment in young adult animals during the period of sexual maturation. Several of these periparturient exposure studies suggest that the observed responses differ from those manifested following gestational DEHP exposure. In an effort to evaluate the consistency of these observations, data were collected from studies analyzing the effects of DEHP on adolescent humans or animals exposed during the periparturient period. This evidence was cumulated by endpoint, the specific timing of DEHP exposure, and the age that the outcome was assessed. The observations were then compared to evidence from exposures during fetal development. The human evidence was difficult to interpret, as exposure timing could not be ascertained. Based on the animal data, several outcomes, including male reproductive system toxicity, showed evidence of differential effects following periparturient exposure. Mechanisms that may explain these differences, such as proposed effects on postnatal androgen biosynthesis, were considered in light of biological plausibility to inform the interpretation of these data. Overall, our analysis suggests that the periparturient period may be a stage of development sensitive to DEHP exposure; however, more comprehensive studies are needed to determine the impact of DEHP exposure on sexually immature adolescents.
Disclaimer: The views expressed are those of the author and they do not represent U.S. EPA policy or guidance.

1006 Diet Acclimation of NZW Rabbits for Use in Embryo-Fetal Development Studies.
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Poor food intake in pregnant rabbits can lead to adverse outcomes including abortion. Initial validation studies with a line of New Zealand White (NZW) rabbits, showed a high incidence of inappetence and adverse pregnancy outcomes. Several studies were conducted to optimize diet acclimation procedures at the breeding facility to match the certified diet used at the Testing Facility. Once the diet acclimation procedure was optimized, a study was conducted to meet the ICH guideline. Poor food intake in pregnant rabbits can lead to adverse outcomes including abortion. Initial validation studies with a line of New Zealand White (NZW) rabbits, showed a high incidence of inappetence and adverse pregnancy outcomes. Several studies were conducted to optimize diet acclimation procedures at the breeding facility to match the certified diet used at the Testing Facility. Once the diet acclimation procedure was optimized, a study was conducted to meet the ICH guideline. The observed dysmorphogenesis of BAs and the impaired migration of the NCCs were found. Treatment-related increase in dysmorphogenesis, no alterations of BA, and no disturbance of NCC distribution in the craniofacial region. In the in vivo study, embryos were harvested on GD 11 from dams gavaged since GD6 with 0, 50, 100, and 180 mg/kg bw/d (the maximal tolerated dose which also causes CP). EPX caused maternal toxicity at 100 and 180 mg/kg; however, these embryos (embryonic tissue conc. = 6.4 mg/l) and dysmophogeneses in 92% of embryos at 30 mg/l (embryonic tissue conc. = 8 mg/l). Embryos exposed to at least 10 mg/ml exhibited an abnormal NCC distribution in the craniofacial region. In the in vivo study, embryos were harvested on GD 11 from dams gavaged since GD6 with 0, 50, 100, and 180 mg/kg bw/d (the maximal tolerated dose which also causes CP). EPX caused maternal toxicity at 100 and 180 mg/kg; however, these embryos (embryonic tissue conc. = 6.4 mg/l) developed similarly to controls. No treatment-related increase in dysmorphogenesis, no alterations of BA, and no disturbance of NCC distribution were found. The observed dysmorphogenesis of BAs and the impaired migration of the NCCs in the in vivo study supports the thesis that EPX alters CYP26-mediated RA metabolism, causing CP. But, despite comparable embryonic tissue levels in the in vivo study, neither the morphology nor the NCCs distribution was affected by EPX. These data do not support the theory that EPX-induced cleft palate in vivo is caused by alterations in RA metabolism.

1008 Embryotoxic Potential of Epoxiconazole In Vitro and In Vivo.
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Some triazoles cause CYP26 inhibition of all-trans retinoic acid (RA) metabolism during organogenesis known to cause cleft palate (CP), via a mode of action (MoA) where altered tissue RA concentrations disrupt neural crest cell (NCC) migration, impairing development of the branching arches (BA). Since very high-dose epoxiconazole (EPX) treatment induces clear maternal toxicity, causing CPs, we examined whether these are also caused by this MoA. Both whole embryo culture tests (WEC) and in vivo studies were performed where the development of the embryos was examined morphologically. NCCs were visualized using whole immunostaining (WIS) for cellular retinoic acid binding protein, and embryonic tissue levels of EPX were analyzed with UPLC. In WEC, embryos were exposed to 0, 1, 3, 10, 20, 30 mg/l for 48 hours starting at gestation day (GD) 9.5. EPX caused alterations in BA in 30% of the embryos at the LOAEC (3 mg/l) and dysmophogeneses in 92% of embryos at 30 mg/l (embryonic tissue conc. = 8 mg/l). Embryos exposed to at least 10 mg/ml exhibited an abnormal NCC distribution in the craniofacial region. In the in vivo study, embryos were harvested on GD 11 from dams gavaged since GD6 with 0, 50, 100, and 180 mg/kg bw/d (the maximal tolerated dose which also causes CP). EPX caused maternal toxicity at 100 and 180 mg/kg; however, these embryos (embryonic tissue conc. = 6.4 mg/l) developed similarly to controls. No treatment-related increase in dysmorphogenesis, no alterations of BA, and no disturbance of NCC distribution were found. The observed dysmorphogenesis of BAs and the impaired migration of the NCCs in the in vitro study supports the thesis that EPX alters CYP26-mediated RA metabolism, causing CP. But, despite comparable embryonic tissue levels in the in vivo study, neither the morphology nor the NCCs distribution was affected by EPX. These data do not support the theory that EPX-induced cleft palate in vivo is caused by alterations in RA metabolism.

1009 Perfluorooctanoic Acid-Induced Cytotoxicity in Primary Cardiomyocyte Culture.
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Perfluorooctanoic acid (PFOA) is a perfluorinated compound (PFC) that is widely used as a polymerization aid in the production of fluoropolymers. PFOA is environmentally persistent and is now detected ubiquitously in biota. Numerous studies have shown that PFOA can induce developmental toxicity in laboratory animal models. Epidemiological studies have demonstrated that PFOA exposure is associated with increased cholesterol levels and cardiovascular disease risk. We have previously reported that PFOA induces developmental cardiotoxicity in chicken embryos and hatchlings. To investigate the mechanism, we developed primary
cardiomyocyte cultures from D10 chicken embryo hearts and evaluated cell viability and reactive oxygen species (ROS) generation after in vitro PFOA exposure. Primary cardiomyocytes were treated with vehicle (0.1% DMSO in medium) or 0.1, 1, 10, 50, 75 or 100 ug/ml of PFOA for 1 or 36h. No statistical differences were detected between untreated and vehicle-treated groups. Viability was decreased by 74.5% in cells treated with 100 ug/ml of PFOA relative to the vehicle group (n=4, P<0.05) at the 1h time point. At the 36h time point, viability was statistically decreased at 10, 50, 75 and 100ug/ml concentrations, relative to the vehicle group (18.1%, 18.7%, 34.6% and 70.0%, respectively; n=4-6, P<0.05). At the 1h time point, an increase in ROS generation was observed at all doses; however, only 50 ug/ml of PFOA statistically increased ROS generation relative to the vehicle group (316.8%; n=3, P<0.05). Our results indicate that direct PFOA exposure to primary cardiomyocytes can induce cytotoxicity and ROS generation. Although additional studies are necessary to verify this effect in vivo, induction of cell death and generation of ROS may partially contribute to developmental cardiotoxicity associated with in ovo PFOA exposure in an avian model.

1010 Estradiol Modulates Paraoxonase-2 Expression in Mouse Brain.

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Paraoxonase 2 (PON2), a member of the PON gene family, is expressed in mouse brain; levels are highest in dopaminergic areas (e.g. striatum), and are higher in astrocytes than in neurons or microglia. As in other tissues, PON2 exerts a potent antioxidant effect in the CNS, and protects mouse neurons and astrocytes against oxidative stress. In all mouse tissues, including the brain, PON2 levels are higher in female than in male mice. In primary striatal astrocytes and neurons PON2 protein level in females is 2-3-fold higher than in males. Levels of PON2 mRNA and lactonase activity show similar gender differences. Male astrocytes and neurons are more sensitive (by 3-4-fold) than female cells to oxidative stress-induced toxicity, though GSH levels do not differ between cells isolated from the two genders. In contrast, no significant gender difference in susceptibility is seen in cells from PON2-/- mice, suggesting that PON2 is a major determinant of gender differences in susceptibility to oxidative-stress-mediated neurotoxicity. Estradiol induces a time-and-concentration-dependent increase in the levels of PON2 protein and mRNA in male (4.5-fold), and also in female (1.8-fold), astrocytes. Such effect is time-and concentration-dependent increase in the levels of PON2 protein and PON2-/- mice, suggesting that PON2 is a major determinant of gender differences although GSH levels do not differ between cells isolated from the two genders. In male astrocytes and neurons PON2 protein and antioxidant effect in the CNS, and protects mouse neurons and astrocytes against oxidative stress associated with in ovo PFOA exposure in an avian model.

1011 Suppression of H19 Methylation in Mouse Exposed to Chlorpyrifos-Methyl during Organogenesis Period.

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Chlorpyrifos-methyl (CPM) is widely used organophosphorus insecticide. In Korea, more than sixty-five tons per year are used in agriculture. In our previous study, CPM showed anti-androgenic endocrine disruption activity. Loss of EDC shows the alteration of methylaion in imprinting gene, which inherit to next generation. This study was performed to examine the effect of CPM on the H19 methylation in mouse. After mating CAST/Ei (C) and B6 (Q), CPM was administered at dose of 4 (CPM4), 20 (CPM20) and 100 (CPM100) mg/kg bw/day from embryonic day (ED) 7 to ED 17. Anogenital distance (AGD) was measured at post natal day (PND) 21. Clinical signs, body weights, feed and water consumption, organs weights, serum hormone values and H19 methylation level of organ and sperm were measured at PND6.

Body weights were significantly (p value <0.01) lower than control until PND 6. AGD was significantly (p value <0.0.0.1) decreased at CPM100 group in male and increased at CPM20 group in female. The weight of thymus and epididymis were significantly (p value <0.0.1) increased at all of CPM treatment in male. In CPM20 group, liver, kidney, heart, lung, spleen, prostate gland and testes were statistically increased. Testosterone level in serum was significantly (p value <0.0.1) increased by CPM treatment in both male and female. H19 methylation level of liver and thymus showed decreased pattern by dose-dependent manner in male. The levels of H19 methylation in sperm were 73.7±6.7% (16% (CTL), 57.8±4.12% (CPM4), 64.2±4.3% (CPM20) and 65.2±4.3% (CPM100). CPM can disturb the early development of offspring development and disrupt H19 methylation in organ and sperm. Those altered methylation pattern may pass down to next generation through sperm.

1012 In Utero Growth Restriction and Dibutyl Phthalate Exposure May Cooperatively Disrupt Steroidogenesis.

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Masculinization of the male reproductive tract occurs in human gestational weeks 10-22, and rat gestational days 16-21. Fetal testes produce a surge of testosterone causing reproductive masculinization. Disruption leads to reproductive abnormalities including cryptorchidism and hypospadias. In humans, in utero growth restriction (IUGR) is a risk factor for cryptorchidism and hypospadias. Rat work shows that fetal steroidogenesis is disrupted by exposure to the anti-androgen dibutyl phthalate (DBP). While these fetal circumstances increase the prevalence of the same birth defects individually, we examined a possible cooperative effect of these treatments on steroidogenesis. A maternal under-nutrition model was examined in the rat. Pregnant Wistar rats were restricted to 50% (n=9) of the food eaten by ad libitum controls (n=8) beginning on gestational day (GD) 3. Restriction through GD17 caused a significant decrease in pup weight with no decrease in litter size. Fetal testes showed decreased testosterone production (40%) by radioimmunoassay. mRNA levels of steroidogenic genes including Scarb1 and Star were significantly decreased in the fetal testis after growth restriction. Food restriction (n=11) from GD3-18 caused no malformations at postnatal days 1, 14, or 28 as compared to controls (n=10). A coexposure study of IUGR and maternal dosing with the anti-androgen DBP was performed examining a cooperative effect on steroidogenesis. Analysis of fetal testes (GD17) following 50% food restriction from GD3-17 (n=11) or exposure to DBP (250mg/kg/day) from GD15-17 (n=7) showed a significant decrease in the expression of the steroidogenic genes Cyp11a1, Cyp17a1, Scarb1, and Star as compared to controls (n=7). When 50% restriction was coupled with DBP exposure (n=10), further decreased steroidogenic gene expression. While IUGR and DBP lead to a decrease in steroidogenesis separately, they also appear to work cooperatively to cause further disruption.

1013 A Comparison of Social Housing Opportunities during Developmental Toxicity Evaluation in Cynomolgus Monkeys.

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The demand for developmental toxicity studies in nonhuman primates (NHPs) has increased, mainly driven by biopharmaceutical drug development. Social housing of NHPs has become mandatory in many countries and this poses a specific challenge since NHPs with timed pregnancies cannot be obtained commercially. The cynomolgus fertility rate in a mating programme is typically 35-45% per mating cycle and 60% per female animal upon repeated mating trials. Hence, social housing of pregnant cynomolgus monkeys requires special approaches until a cage is populated by animals with confirmed pregnancy. Formation of new groups of sex-matched animals is challenging due to the extent that some animals exhibit transient amenorrhea. Concern has been raised that social housing of pregnant animals might interfere with maintenance of gestation due to rank order fights, incompatibility, and other group related adverse effects. On the other hand, in a social setting, animals experience birth events and handling of neonates by watching cage mates, and infants have cage mates of comparable ages to interact with. In the present work, qualitative assessment in more than 200 hundred pregnant control animals under social housing indicated good compatibility with little aggression toward each other. Interestingly, a reduction of pre- and postnatal loss by 25-30% was encountered under social housing. Conceivably, the delivery process and management of the newborn was facilitated by social maternal housing. Moreover, maternal and infant body weight gains were significantly increased under social housing.
Feasibility of Mouse Continuous Intravenous Infusion for Fertility and Embryo-Fetal Development Studies.


The rat and rabbit are routinely used in pre-clinical reproductive and developmental toxicity (DART) studies. However, where the rat and/or rabbit are not suitable, the mouse is an alternative model. A model of continuous intravenous infusion using a surgically-implanted femoral vein catheter with tail cuff exteriorisation has been previously developed in the mouse. The reliability of this method decreased over time, with a success rate of only 74% after 28 days because of tail cuff constraints due to animal growth. This study aimed to (1) determine whether a larger diameter tail cuff (6 mm) could extend the continuous infusion duration, and (2) determine the feasibility of evaluating DART parameters in mice using this surgical model in a combined fertility and embryo-fetal development study.

33 CD-1 female mice (approximately 7–9 weeks old) were surgically implanted with a femoral vein catheter exteriorised with a 6 mm tail cuff. After 6 days of recovery, they were intravenously administered 0.9% sodium chloride, by continuous infusion (4 mL/kg/hr) for 2 weeks prior to pairing, during pairing (with un-catherised males) and through Gestation Day (GD) 18 (42 days). On GD18, females were killed and uterine contents examined. Technical failure occurred in 9 females; this was attributed to the larger diameter of the tail cuff, which slipped down and allowed the catheter to be chewed. Of the 24 female mice remaining on study at the end of pairing, 23 (96%) were successfully mated and 22 (96%) were confirmed pregnant. Cather failure or poor tail conditions resulted in early removal of 5 additional mice. However, those surviving to necropsy, uterine and foetal data were within background ranges, indicating no effects on reproductive or developmental parameters. It is concluded that continuous intravenous infusion via this surgical model is a viable method of dosing mice in DART studies.

Addressing the Mode of Action for Developmental PFOA-Induced Mammary Gland Delays.

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Perfluorooctanoic acid (PFOA) exposure alters mammary gland (MG) development and function in mice. The mode of action (MOA) for these effects has not been elucidated, although activation of peroxisome proliferator activated receptor alpha (PPARα) has been implicated. To begin elucidation of a MOA, time pregnant CD-1 mice were gavaged with 0.0, 0.01, 0.1, or 1.0 mg PFOA/kg body weight on gestation days (GD) 10–17. Female offspring MGs were removed and prepared for RNA on multiple postnatal days (PND). A comparison of MG RNA from control and 1.0 mg/kg groups with Affymetrix 420.2 microarrays found different regulation of 710 and 41 genes at PND 7 and 14, respectively. Adiponectin, estrogen receptor alpha, and WNT expression were reduced in PFOA-treated glands. PPARα expression was not changed. These data were validated by PCR of MG RNA, using TBP as a reference gene. At PND 7, PFOA induced differential regulation of several genes compared to controls, even at 0.01 mg/kg. When considering developmental scores, glands with higher scores or more normal growth had increased RNA expression of WiFi1, ESR1 and PGR at PND 21 and FABP3 and UCP1 at PND 56. PND 56 MG sections stained for PPARα protein displayed robust expression in all glands regardless of treatment, yet there was higher expression in stromal tissues surrounding ducts in PFOA exposed mice compared to controls. Collectively, these findings suggest that PPAR pathway genes are altered in conjunction with PFOA induced MG changes along with other pathways. Observed altered genes are involved in signaling of all PPAR isoforms including γ, δ, and β/δ. Both up and down regulation of these genes were found which suggests that multiple PPAR isoforms may work together to regulate PFOA-induced MG aberrations. In addition, prenatal PFOA exposure at 0.01 mg/kg leads to mammary gene changes at human-relevant levels. Future investigations will examine related gene pathways and protein levels compared to observed morphological and hormonal outcomes. This abstract does not necessarily reflect NIEHS policy.

Evaluation of Reproductive and Developmental Toxicity of Multiwall Carbon Nanotubes in Pregnant Mice After Intratracheal Instillation.

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Some studies have reported that maternal exposure to nanoparticles may induce teratogenicity or transfer to their fetuses and affect the development and function of the central nervous systems. In order to evaluate the reproductive and developmental toxicity of multi-wall carbon nanotubes (MWCNTs) via inhalation exposure, we conducted intratracheal instillation study of MWCNTs dispersed in two different media (blood serum obtained from female mice and 2% of sodium carboxymethyl cellulose (CMC-Na) solution) in pregnant mice. MWCNTs (MWN7-T) dispersions were prepared by ultrasonication using an ultrasonic bath for 30 min. The above MWCNT dispersions were administered to pregnant C57BL/6JICR mice with gestation day 9 at dosage of 0.3, 3.0, and 5.0 mg/kg bw. Five mice per group were evaluated. The mice were dissected at 18 days of gestation period. MWCNT dispositions in the tissues and the effect to embryos and fetuses were evaluated. The MWCNTs were deposited in the lungs as the black spots. No statistically significant difference between the control group and MWCNT-exposed groups in the number of corpora lutea, implantations, dead fetuses (early or late resorption site), live fetuses, and sex ratio. However, body weight of fetuses and placental weights were dose-dependently decreased in the both MWCNT-exposed groups. Furthermore, external malformations (i.e., oligodactyly; extensivecontracture, cleft lip) were observed in the 3 mg/kg of CMC-Na treated MWCNT-exposed group and 5 mg/kg of blood serum treated MWCNT-exposed groups. Further information is needed to clarify the potential for the reproductive and developmental toxicity of MWCNTs.

Histology of Selected Organs from the Göttingen Minipig Fetuses from Days 60 (Midterm) and 110 (Term) of Gestation.


In a previous study, we demonstrated that chemically induced fetal abnormalities could be detected in the Göttingen Minipig shortly after mid term. At this time, it is possible to sex the minipig fetus by observation of the external genital area, although at gross soft tissue examination, the location, size and shape of the male and female gonads are similar at this stage. The aims of the present study were to compare the histological features of selected organs in the Göttingen Minipig fetus at GD 60 and GD 110 and to validate histologically the method of sexing the fetuses by macroscopic observation of the external genital area.

Four 7- to 13-month-old virgin Göttingen Mining females were mated. Caesarean sections were performed on two females at mid term (GD 60) and on two others at term (GD 110). The fetuses were weighed, sexed and submitted to standard gross external and visceral examinations. Selected organs (adrenal glands, heart, kidneys, liver, lungs, ovaries, thymus, thyroid gland, urinary bladder, uterus, vagina, epididymis and testes) were sampled and fixed in 10% formalin or modified Davidson’s fluid. Male five and 5 female fetuses obtained at GD 60, and 2 male and 6 female fetuses obtained at GD110 were available for histological examination. The microscopic appearances and the differences between the two timepoints were described for each organ.

The histological appearance of the organs at GD110 was similar to that of juvenile minipigs. At GD 60, however, although all organs were identifiable, maturation and growth were obviously still incomplete. This study reported the histology of selected organs in minipig fetuses at GD 60 and GD 110. At GD 60, there was complete concordance in establishing the sex of each fetus by microscopic examination and evaluation of the external genitalia.

Mono-Ethylhexyl Phthalate (MEHP) Exposure Reduces Embryonic Nutrition and Induces Structural Defects in Mouse Whole Embryo Culture.

K. E. Sant and C. Harris. *Environmental Health Sciences, University of Michigan, Ann Arbor, MI.*

Di-2-ethylhexyl phthalate (DEHP) is a highly lipophilic endocrine disruptor and is the most abundant phthalate plasticizer in the environment. Exposure to DEHP and its primary active metabolite, mono-ethylhexyl phthalate (MEHP), have been associated with several adverse health effects, ranging from obesity to reproductive
and M. Harrass.

Di-n-butyl phthalate (DBP) is a common plasticizer used in a variety of consumer products. Its major metabolite is its monoester. Exposure to DBP is widespread, and its potential toxicity has been, and continues to be, investigated. The objective of this work was to develop, validate, and apply a method to quantify mono-n-di-n-butyl phthalate (MBP), the major metabolite of DBP, in rat plasma and amniotic fluid.

The method is being applied for the analysis of MBP in plasma and amniotic fluid samples from rats administered 0, 300, 1000, 3000, or 10,000 ppm DBP in feed.

Developmental and Reproductive Studies in Sprague-Dawley Rats with Gevkizumab, a Monoclonal Antibody Targeting IL-1 Beta.

1020

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Gevkizumab, also coded XOMA 052 or 578/989, is a humanized IgG2 monoclonal antibody that binds with high affinity to human interleukin-1 (IL-1) beta. Gevkizumab inhibits the activation of the IL-1 receptor, thereby modulating the cellular signaling events producing inflammation. Currently, gevkizumab is in Phase 3 clinical trials for non-infectious uveitis and Behcet’s uveitis and in Phase 2 proof-of-concept studies for moderate to severe acne and erosive osteoarthritis of the hands. Developmental and reproductive studies (pre/postnatal development) in Sprague Dawley rats evaluated the effects of gevkizumab when administered subcutaneously once weekly at 30, 300, and 90 mg/kg/dose. The rat (usual rodent model for evaluating developmental and reproductive toxicity) was selected due to similar binding affinity and in vitro functional activity between human and rat IL-1 beta. Standard ICH S5(R2) study designs were followed. Observations included clinical signs, body weight, food consumption, drug levels in serum and milk, anti-drug antibodies (ADA), maternal uterine and ovarian examinations, fetal evaluations, macroscopic pathology, reproductive performance, parturition, lactation to weaning, and F1 physical growth and development. Gevkizumab did not produce any mortality or significant adverse effects. Measurement of gevkizumab in serum (maternal, fetal, neonate) and maternal milk confirmed adequate exposure during the dosing periods. ADAs were detected in a few dams, without neutralizing activities. The NOAEL for maternal and developmental toxicity including teratogenicity, general reproductive toxicity in male and female rats, and general toxicity in the dam, including developmental and reproductive toxicity in the F1 offspring was considered to be at least 90 mg/kg/dose, the highest dose level tested. No gevkizumab-related findings were observed in any of the reproductive and developmental studies.

1021 Weight of Evidence Considerations for Developmental Toxicity Classification of Boric Acid. W. Ball and M. Harrass. Rio Tinto Minerals, Greenwood Village, CO.

Although reproductive and developmental effects have been demonstrated in laboratory animals exposed to high doses of boric acid (BA), similar effects have not been observed in highly exposed human populations or workers. Workers in boron (B) mining/processing industries represent the maximum possible human exposure. A weight of evidence approach was used in evaluating numerous independent studies on the determination of the hazard of BA to humans. Information that was considered together included results of in vitro tests, animal data, worker exposure data, epidemiological studies and mechanistic data. Extensive evaluations of sperm parameters in highly exposed workers in Turkey and China demonstrated no effects on male fertility. No evidence of developmental effects in humans attributable to B has been observed in studies of populations with high exposures to B. Collectively, the epidemiological studies consistently show an absence of effects in highly exposed populations. A comparison of blood, semen and target organ B levels in studies of lab animals and B workers shows B industry worker exposures are lower than untreated control rats. Recent studies provide evidence that BA may act by similar mechanisms in causing developmental effects in mice as sodium salicylate (natural deacetylated form of aspirin) including effects on Hox gene expression and inhibition of embryonic histone deacetylases. Although aspirin is known to cause developmental effects in laboratory animals, similar effects have not been seen in controlled human studies. Similar mechanisms of action of BA and aspirin and the absence of effects in humans ingesting aspirin suggest that BA related developmental effects in humans are unlikely. Also, there is evidence that Zinc (Zn) interacts with B in the body reducing the toxicity of B. Zn levels in soft tissue in humans are over 2 x greater than in lab animals explaining in part the absence of fertility and developmental effects in humans. Based on the total weight of evidence, the data show that it is improbable that BA will cause reproductive or developmental effects in humans.

1022 Embryonic DNA Repair and Ethanol-Initiated Behavioural Deficits in Osoguanine Glycosylase 1 (OGG1) Knockout Mice: A Role for Oxidatively-Damaged DNA and Protection by a Free Radical Spinning Agent. L. Miller1, D. J. Pinto1 and P. G. Wells1, 1Pharmacology and Toxicology, University of Toronto, Toronto, ON, Canada; 2Pharmaceutical Sciences, University of Toronto, Toronto, ON, Canada.

A mother’s consumption of alcohol (ethanol, EtOH) during pregnancy can cause a spectrum of structural, cognitive and behavioural problems in the developing child termed Fetal Alcohol Spectrum Disorders (FASD). Reactive oxygen species (ROS) have been implicated in the mechanism of behavioural teratogenesis, although the role of embryonic DNA damage and repair are unclear. To determine the latter, DNA repair-deficient heterozygous (+/-) osoguanine glycosylase 1 (OGG1) knockout mice were mated, and pregnant dams were treated once on gestational day 17 with EtOH (2 g/kg, i.p.) or its solvent vehicle, with or without pretreatment with the free radical spin trapping agent alpha-phenyl-N-tert-butyl nitronitrone (PBN) (40 mg/kg, i.p.). Saline-exposed progeny exhibited an oggl gene dose-dependent learning deficit in the passive avoidance test compared to wild-type (WT) littermates, demonstrating for the first time a phenotype in OGG1-deficient mice. EtOH-exposed progeny exhibited an enhanced learning deficit compared to saline controls at 6 and 9 weeks of age, also in an oggl gene dose-dependent fashion. PBN pretreatment significantly protected both WT and KO progeny, although this protection for EtOH was slightly less effective in +/- and -/- KO progeny. These results provide the first evidence to date that ROS-initiated embryonic DNA oxidation is involved in EtOH-initiated behavioural deficits, and embryonic DNA repair status may be a determinant of teratological risk. (Support: Canadian Institutes of Health Research)
Polychlorinated biphenyls (PCBs) are persistent environmental pollutants that may cause adverse health effects. Previous in vitro studies including ours have shown that various PCB congeners affect neurodevelopment through different mechanisms. However, behavioral alterations induced by lactational exposure to PCB and its neurochemical mechanisms have not yet been fully studied. In the present study, hydroxylated-PCB 106 (OH-PCB 106; 4-hydroxy-2,3,3',4,5'-pentachlorobiphenyl) was administered to lactating rat and mouse dams via gavage every second day from day 3 to 13 after delivery. The exposure did not affect the body weight of the dams or the physical development of the newborn pups in both sexes. Male rats and mice exposed to OH-PCB 106 showed hyperactivity that was characterized by increased locomotor activity in open field and circadian period. OH-PCB 106-exposed rats displayed abnormally high levels of dopamine and D2 dopamine receptor (D2DR), but not D1DR and D5DR, in the striatum, an important center for the coordination of behavior. These findings indicate that OH-PCB 106 has a significant neurotoxic effect on rodent behavior, which may be associated with increased D2DR mediated signals.

### Di-isobutyl Phthalate (DIBP) Hazard Identification


The hazard potential for DIBP is being evaluated as part of EPA’s Integrated Risk Information System (IRIS) Toxicological Review. DIBP is a plastifier that confers flexibility and durability in industrial and consumer products. The epidemiology and animal toxicology databases are both relatively small. The small DIBP epidemiological database includes studies that assessed the relationship between urinary concentrations of the DIBP metabolite mono-isobutyl phthalate (MIBP) and developmental, neurodevelopmental, immunological or breast cancer outcomes. There is limited support for associations between MIBP and inflammatory biomarker levels, and decreased masculine play behavior. The animal toxicological database includes studies that assessed male reproductive developmental endpoints that constitute the phthalate syndrome after in utero DIBP exposure. The largest developmental study, Saillenfait et al. (2008), reported changes in anogenital distance, male reproductive organ weights, and litter incidence of phallic syndrome endpoints in the lower dose range after early gestational exposure. Other studies observed increased fetal mortality, male postnatal and adult growth decrements, decreased fetal testicular testosterone and changes in expression of genes involved in androgen production pathways. The developmental reproductive effects observed in animal studies are consistent with the reduced testicular testosterone mode of action that is well-characterized for developmentally toxic phthalates. Effects on liver and kidney weight and function in both male and female adults were also reported. Research needs include epidemiologic studies that examine DIBP exposure, testosterone and related outcomes throughout development as well as multigenerational reproductive toxicity studies and cancer bioassays. The views expressed are those of the authors and do not necessarily reflect the views or policies of the US EPA.

### The Devtox Project: A Comprehensive Source of Information on Developmental Abnormalities

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The potential of a compound to cause adverse effects in the developing embryo or fetus is an important consideration in human health risk assessment. The terms and diagnostic criteria used to describe fetal anomalies in animal experimentation need to be consistent from one laboratory to another and must be harmonized between regulatory agencies. Consequently, the DevTox Project (www.devtox.org) has the main objectives to harmonize the nomenclature used to describe developmental anomalies in laboratory animals and to assist in the visual recognition of developmental anomalies with the aid of photographs. The use of a harmonized and internationally accepted nomenclature is a basic requirement for the operation of DevTox. A first approach of establishing such a harmonized terminology was made in 1997 by a publication of the International Federation of Teratology Societies IFTS (Teratology 55; 249-292, 1997). This terminology has recently been updated (Reproductive Toxicology 28, 371-434, 2009). In addition, a series of Berlin Workshops working definitions for the two classification categories “malformation” and “variation” were agreed. The results of all these activities were used to establish the DevTox database. The easy-to-use Web interface allows different views of the nomenclature, images and data and a quick navigation throughout the complete site. DevTox currently contains more than 2,500 images, showing examples of external, skeletal, soft tissue and maternal-fetal anomalies in rats, mice, rabbits, hamsters, primates, Guinea...
pigs, minipigs, dogs and birds. It provides short descriptions of each finding and in some cases synonyms and further diagnostic notes as well as a hierarchical structure for the localizations. The system is publicly accessible and allows the electronic download of the current version of the harmonized terminology.

1028 Reference Images for the Skeletal Examination of Cynomolgus Monkey Fetuses and Thalidomide-Induced Malformations.
O. Foulon, F. Specia, C. Dauzat, M. Da Silva, P. Barrow, B. Pulate and R. Forster. GToxLAB, Evreux, France.

In this poster selected images are presented to illustrate the normal and (drug-induced) abnormal skeletal morphology of the cynomolgus monkey at 100 days post-coitum. Very limited information is currently available in the literature in terms of reference images for the skeletal examination of cynomolgus monkey (Macaca fascicularis) fetuses in teratology studies. This poster will present photographs of the fetal skeleton on day 100 post-coitum. Normal (control) fetuses will be presented and compared with fetuses showing thalidomide-induced malformations. A total of 20 control and 3 thalidomide-treated fetal specimens were photographed. Dams were 3 to 8 years old at mating and weighed 2.0 to 4.5 kg. Mating was confirmed by the presence of sperm in a vaginal smear during cohabitation with a male. Pregnancy was confirmed 18 or 19 days later by ultrasound examination. The treated dams were given 20 mg/kg/day of thalidomide by oral gavage from days 25 to 30 of gestation. The control dams received a similar volume of water (10 mL/kg). The fetuses were removed by caesarean section on day 100 of gestation. Fetal and placental weights were recorded. Morphometric measurements (head circumference and crown-rump-length) were taken. Each fetus was then given a detailed necropsy examination prior to processing for skeletal examination by the Dawson method: the ossified skeletal structures were visualised following clearing of the soft tissues in potassium hydroxide and staining with Alizarin Red S. The normal variation in the normalisation of the number of sows mated per day according to the facility capacity to perform caesarean sections on the required day of gestation. The data presented show that our methods give a mating success and an average litter size that are suitable (e.g. due to issues of metabolism) a non-rodent alternative choice of species is available (e.g. due to issues of metabolism) a non-rodent alternative choice of species is available (e.g. due to issues of metabolism) a non-rodent alternative choice of species is available (e.g. due to issues of metabolism) a non-rodent alternative choice of species is available.
1 and Rho kinase inhibitor (HA1077). The response to phenylephrine was studied in the presence and absence of COX inhibitor, indomethacin. Dose-response curves, maximum stress and EC50 values were different for silver NP exposed animals when compared with vehicle controls. Reciprocal changes were seen between the aortic and uterine vessels: greater vasoconstriction in uterine artery and greater vasodilation in aorta following 110 nm silver exposure. Exposure to 20 nm silver NP increased the HA1077-mediated vasorelaxation in aortic vessels, suggesting a possible mechanism underlying the changes of agonist mediated vasconstrictor response following NP exposure. NP suspensions in citrate lead to higher stress generation in both vessels. IV exposure to silver NP during pregnancy induces particle size and vehicle dependent changes in vascular reactivity which can potentially influence blood supply to the fetus. This work is supported by NIEHS U19 ES019525.

Flubendazole confirmed to be embryolethal and teratogenic through its cytostatic properties. At 6.32 mg/kg/day it initially induced a block of embryonic development through its cytostatic action, followed by cytotoxicity which led to 100% embryolethality within 48h while at 3.46 mg/kg/day, it markedly reduced embryonic development through its cytostatic action but did not show cytotoxic effects. Therefore, embryos still alive markedly retarded with consequent morphological alterations in specific areas namely brain vesicles, eye, maxillary structures and body axis.

In a previous study, we have demonstrated that fetal abnormalities associated with a known human and swine teratogen (Pyrimethamine) could be detected in the minipig with fetal examinations performed close to mid-term instead of at term. The principal objective of the present study was to validate a double staining method for bone and cartilage of minipig fetuses obtained by caesarean section close to gestation day (GD) 60 in order to support skeletal examinations.

Two virgin Göttingen Minipig females were mated at the testing facility and subsequently submitted to caesarean section close to GD 60. Ten fetuses (5 in each litter) were available for the trials and the sexes were evenly split.

The fetuses were used in a number of staining trials based on our own established methods for other species, including the rat and rabbit. In the first step, the fetuses were stained, without prior fixing, following evisceration and skinning. Varying concentrations of ethanol, mixed together with Alizarin red S, Alcian blue and acetic acid, were tested to obtain the desired results. In order to establish the best conditions to fix the staining, the fetuses were then transferred to baths containing different concentrations of ethanol for varying durations. Following fixation, the specimens were transferred into potassium hydroxide solutions for maceration of the soft tissue. Finally, the soft tissues were cleared, firstly using Mals solution followed by glycerol at increasing concentrations. The fetuses from the successful trial were then ready for skeletal examination.

The technique is slightly more labour intensive than the single staining technique using Alizarin red S only. However, the time required to process the specimens for examination is approximately the same for both methods. Moreover, this new technique enhances the examination process for mid-term minipig fetal skeletons, at a time when ossification of the skeleton is obviously less advanced than at term. Our next step is to generate an atlas of the mid-term minipig fetal skeleton.

BMS-708163 (avagacestat) inhibits γ-secretase (GSI), a protease that cleaves amyloid precursor protein to produce amyloid β (Aβ) and amyloid plaques that are prominent lesions in Alzheimer’s disease (AD). As part of the nonclinical reproductive safety evaluation, studies of embryofetal development as well as fertility and early embryonic development were conducted in female Sprague-Dawley rats. In the embryo-fetal development study, BMS-708163 was administered (oral gavage) to pregnant rats at 3, 10 and 30 mg/kg/day from Gestation Day (GD) 6 through GD 15. Cesarean sections were conducted on GD 21. Assessments of external well-being included clinical observations, body weight, and food consumption; evaluation of fetal viability, morphology and body weights comprised the assessment of developmental toxicity. BMS-708163 was a selective developmental toxicant at all doses tested, with dose-related increases in the incidence and severity of fetal malformations and variations in the absence of maternal toxicity. Malformations and variations included axial and appendicular skeletal anomalies. In the study of fertility and early embryonic development in female rats, BMS-708163 was administered for 14 days prior to cohabitation; then throughout cohabitation at all doses tested, with dose-related increases in the incidence and severity of embryonic toxicity. BMS-708163 was a selective developmental toxicant at all doses tested, with dose-related increases in the incidence and severity of embryonic toxicity.

In order to evaluate the embryotoxic nature of Flubendazole, embryos of dams treated as above, were evaluated on GD 11.5 and 12.5, as a window for observing the origin of alterations detected at term.

Data from a previous whole embryo culture study. Fetuses were evaluated on GD 20. Flubendazole was embryolethal from 6.32 mg/kg/day (Cmax after single administration 0.801 ug/mL). In addition Flubendazole was teratogenic at 4.56 mg/kg/day (Cmax after single administration 0.539 ug/mL). No increase in resorptions was observed when 80% of fetuses showed malformations namely exencephaly, microcephaly, micro/anophthalmia, small/absent kidneys, markedly enlarged renal pelvis/ureters, absent tail, anal atresia and disruption of skeletal ossification of head, spine and thoracic cage.

Flubendazole did not interfere with rat embryonic development, apart from a minimal reduction in fetal and placental weights, at 2 mg/kg/day (Cmax after single administration 0.389 ug/mL) with treatment restricted to two days during pregnancy and with a limited number of animals.

In order to evaluate the embryotoxic nature of Flubendazole, embryos of dams treated as above, were evaluated on GD 11.5 and 12.5, as a window for observing the origin of alterations detected at term.
1037 Influence of Acidic Extraction Conditions on Formazan Assay: Assessment of Test Substances Using In Vitro EpiSkin Skin Irritation Test Method by HPLC and Colorimetry.

The skin irritation potential of a test substance is typically determined by measuring cell viability in treated tissues by means of the colorimetric MTT reduction assay after topical application of a test substance. In the EpiSkin validated skin irritation test method, tetrazolium salt-based formazan assay is performed by using acidic isopropanol extraction conditions (ESAC 2007, OECD TG 439 2009).

The current work evaluated the data obtained with the EpiSkin model for the 20 reference chemicals (listed in the OECD TG 439) following formazan extraction with acidified (IPA) or non-acidified (IP) isopropanol solutions to evaluate the influence of acidic extraction conditions. Therefore the objectives of this work was to establish the use of HPLC measurements as a complementary assay to standard photometry assay for detection of reduced MTT, a known limitation for test substances that are highly coloured.

The cell viability in EpiSkin quantitatively measured after acidified (IPA) or non-acidified (IP) isopropanol extractions from tissues were equivalent ranging from 6.4 to 104% by colorimetry (reference method). Following HPLC measurement, formazan quantitative analyses were comprised between 7.9 to 99.2%. The standard deviation between those 2 conditions was +/- 18%. Therefore, within-laboratory variability assessed in 3 runs showed similar concordance for classification for each condition. The sensitivity (based on 10 GHS Cat.2 substances) was 90% and the specificity (based on 10 No-category substances) was 70%. The HPLC protocol provides data comparable to those using standard photometry assays for 20 test substances indicative of the use of HPLC as a complementary assay to photometry in EpiSkin skin irritation test method. Therefore, those results suggest that acidic isopropanol extraction conditions did not affect the EpiSkin skin irritation outcomes.

1038 Cytotoxic Effects of Dicyclohexylamine and Three Metalworking Fluids on Human Epidermal Keratinocytes.
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Many of the 1.2 million workers in the metal machining industry within the United States will be exposed to metalworking fluids via the dermal route, making it a major occupational health and safety concern. Dicyclohexylamine (DCHA), an anticorrosive agent used in the metalworking industry to prevent corrosion of fabricated materials, is known to permeate the skin and could potentially elicit a toxic response on human epidermal keratinocytes (HEK). HEK were exposed only to DCHA or one of the following three generic metalworking fluids: soluble oil, synthetic oil and semi-synthetic oil for 24h. Cells were dosed with solutions ranging from concentrations of 200-5000 μg/mL. The highest concentration of 5000 μg/mL resulted in an 80-85% and 81-85% decrease in cell viability, respectively, in comparison to the media control. For synthetic oil, there was a 52-μg/ml resulted in an 80-83% and 81-85% decrease in cell viability, in comparison to the media control. The soluble oil and semi-synthetic oil dosed exposed to 200-5000 μg/mL is the relevant occupational exposure dose in the metal machining industry. Cells produced 1.29 μg/mL, determined by Tripal Blue assay. Besides, this concentration of the dye induced cell cycle arrest at G2 phase. Using the clonogenic assay, we observed that 2.5μg/mL of the dye was sufficient to inhibit completely cell growth after 48 hours of exposure. In the artificial skin model, we demonstrated that the exposure to Basic Red 51 at IC50 reduces the skin thickness, decreasing the number of cell layers. We also observed that the cells appear to be injured and undergoing apoptosis. It is important to point out that the IC50 used for these experiments (12.9 μg/mL) is much lower than the commercial concentration (2000μg/mL). We suggest that similar effects could be induced in humans after exposure to the Basic Red 51, mainly considering the effects in artificial skin. So, we concluded that the use of Basic Red 51 for cosmetic purposes should be carefully evaluated.

1040 Cytotoxic Effect of the Temporary Hair Dye Basic Red 51 on Human Keratinocytes (HaCaT): Development and Application on a Reconstructed (3D) Artificial Skin.
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Nowadays, hair dyes are being commonly used for cosmetics purposes. However in 2001, the Scientific Committee on Cosmetic products and Non-Food Products intended for consumers (SCCNFP) of the European Union recommended that hair dye ingredients should be re-analyzed. Since 2006, 22 hair dyes have been banned for not being considered safe. On the present work, we evaluated the cytotoxicity of a temporary hair dye “Basic Red 51” (CAS: 77061-58-6), an azo dye (N-N-) that is currently being revised by the EU. We tested the toxicity of the “Basic Red 51” in immortalized keratinocytes (HaCaT) that are the primary route of exposure. Moreover, we reconstructed a human (3D) epidermis using HaCaT cells and dermal fibroblasts in order to create a new approach that would better represent the real exposure. For HaCaT cells in monolayer, the Basic Red 51 IC50 was 12.9 μg/mL, determined by Tripal Blue assay. Besides, this concentration of the dye induced cell cycle arrest at G2 phase. Using the clonogenic assay, we observed that 2.5μg/mL of the dye was sufficient to inhibit completely cell growth after 48 hours of exposure. In the artificial skin model, we demonstrated that the exposure to Basic Red 51 at IC50 reduces the skin thickness, decreasing the number of cell layers. We also observed that the cells appear to be injured and undergoing apoptosis. It is important to point out that the IC50 used for these experiments (12.9 μg/mL) is much lower than the commercial concentration (2000μg/mL). We suggest that similar effects could be induced in humans after exposure to the Basic Red 51, mainly considering the effects in artificial skin. So, we concluded that the use of Basic Red 51 for cosmetic purposes should be carefully evaluated.

1041 Skin and Eye Lesions in the Long Evans Rat Produced by Oral 8-Methoxy-Proronal (8-MOP) Application and Subsequent UV-A Irradiation.
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For assessment of the dermal and ocular response to the model phototoxican 8-MOP, 6 female HsdBluLE (SPF) Long Evans rats received a single dose of 5.0 mg/kg body weight 8-MOP by oral gavage. 6 control rats were treated with the vehicle only. One pigmented and one unpigmented skin site were clipped free of hair and irradiated with 15 J/cm² UV-A (Bio-Spectra irradiation chamber, Vilber Lourmat Deutschland GmbH). Irradiation was performed 30 min after 8-MOP application under pentobarbital anaesthesia. Visual assessment of the skin 4 h after irradiation showed erythema on the irradiated pigmented and non-pigmented skin of 8-MOP-treated rats and in control rats which increased in severity in 8-MOP-treated rats after 20 h and receded in control rats . Injection of Evans Blue showed moderate vascular leakage 24 h and 48 h after irradiation in all 8-MOP-treated rats, while only few of the control rats were affected at a lower severity. Histopathologically, corneal single cell necrosis was recorded in the irradiated eye of 2 of the 8-MOP-treated rats, but not in control rats. Dermal lesions in pigmented and unpigmented skin of 8-MOP-treated rats consisted of focal/multifocal epidermal hyperplasia, focal erosions/scab formation, ulceration, subcutaneous and/or dermal infiltration, follicular inflammation, focal/multifocal epidermal necrosis

1039 Aryl Hydrocarbon Receptor Repressor (AhRR) Function Revisited: Repression of Cyp1 Activity in Human Skin Fibroblasts Is Not Related to AhRR Expression.
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The skin reacts to environmental noxae by inducing cytochrome P450 (CYP)-catalyzed reactions via activation of the aryl hydrocarbon receptor (AhR). A drawback of this response is the generation of oxidative stress, which is especially dangerous for postreplicative cells such as dermal fibroblasts, in which damage may accumulate over time. Accordingly, in dermal fibroblasts, CYP1 expression is repressed and it has been proposed that this is due to the AhRR, which is supposedly overexpressed in fibroblasts as compared with other skin cells. Here, we revisited this AhRR hypothesis, which has been mainly based on ectopic overexpression studies and correlation analyses of high AhRR gene expression with CYP1A1 repression in certain cell types. In primary human skin fibroblasts (NHDFs) of 25 individuals, we found that (i) the AhRR was expressed only at moderate RNA copy numbers and that, against the common belief (ii) CYP1A1 mRNA expression could be induced by AhR activators. However, even the highest induction did not translate into measurable CYP1 enzyme activity, and neither basal expression nor mRNA inducibility correlated with AhRR expression. In addition, enhancement of CYP1A1 mRNA expression by trichostatin A, which inhibits AhRR recruited histone deacetylases at the CYP1A1 promoter, failed to induce measurable CYP1 activity. Finally, AhRR deficient mouse embryonic fibroblasts were not induced to biologically relevant CYP1 enzyme activity despite impressive mRNA induction. These data clearly indicate that repressed CYP1 activity in NHDFs is not causally related to AhRR expression, which may serve a different, yet unknown, biological function.
13-Week Dermal Toxicity of Triclosan in B6C3F1 Mice.

M. M. Vanlandingham, L. Fang, F. A. Beland and W. A. Harrouk

Triclosan [5-chloro-2-(2, 4-dichlorophenoxy)phenol] is widely used as an antimicrobial agent, and humans in all age groups have the potential to receive lifelong exposures. We recently demonstrated that the dermal application of triclosan to mice results in systemic distribution of the compound. The goal of the present study was to evaluate the toxicities of triclosan administered dermally to B6C3F1 mice for 13 weeks and provide a scientific basis for dose selection for a subsequent chronic dermal carcinogenicity study. Groups of 10 male and 10 female mice received dermal application of 0, 5.8, 12.5, 27, 58, or 125 mg triclosan in ethanol per kg bw daily seven days per week for a period of 13 weeks. All mice survived the 13-week treatment. Body weights of female mice were not affected, while male mice administered 58 and 125 mg/kg weighed 91% and 83% of the control male mice. The mean weight of livers from females receiving 58 and 125 mg/kg was 11% and 40% greater than the controls; in males, the highest dose led to a higher mean liver weight (38% greater than the controls). Skin lesions (epidermal hyperplasia, inflammation, necrosis, ulceration, and parakeratosis; dermal fibrosis and inflammation) were observed in both sexes. The severity of the lesions was minimal at the 12.5 mg/kg dose, and both the severity and incidence of the lesions increased as the dose was increased. The highest dose of triclosan increased the percentage of reticulocytes in both sexes and the percentage of micronucleated normochromic erythrocytes in females; in addition, the 58 mg/kg dose increased the percentage of reticulocytes in females. A significant dose-dependent decrease in the levels of T4 and cholesterol was observed in both sexes. There were no differences between treated and control mice in sperm or vaginal cytology measurements. The NOAEL, under the conditions of this subchronic assay, was 5.8 mg/kg. (Supported by Interagency Agreement between NCTR/FDA IAG #224-12-0003 and NIH AES12013.)

Dermal Toxicokinetic Studies of Triclosan in B6C3F1 Mice.

L. Fang, M. M. Vanlandingham, F. A. Beland and W. A. Harrouk

Triclosan [5-chloro-2-(2, 4-dichlorophenoxy)phenol] is widely used as an antimicrobial agent in personal care products, household items, and clinical settings. Due to its extensive use, there is potential for humans in all age groups to receive lifelong exposures to triclosan; however, data on the chronic dermal toxicity/carcinogenicity of triclosan are still lacking. The goal of the present study was to determine the absorption, distribution, metabolism, and excretion of triclosan following dermal administration to B6C3F1 mice. A single dose of [14C(U)]triclosan (10 or 100 mg per kg bw, 10 μCi per mouse) was administered to mice in two separate experiments: a vehicle selection experiment to compare the bioavailability of triclosan in male mice using propylene glycol, ethanol, and a generic lotion as vehicles; and a toxicokinetic experiment to evaluate sex differences in the absorption and deposition of triclosan and its metabolites. Three mice per sex in each group were killed at 0.5, 1, 2, 4, 6, 8, 12, 24, 48, and 72 h after administration, and excreta and tissues were collected and analyzed for radioactivity. Ethanol has the best properties of the vehicles evaluated. Maximum absorption of triclosan (80 - 86%) was obtained at 12 h after dermal application. There was some saturation of absorption at the 100 mg/kg dose. Radioactivity appeared in the excreta and in all tissues examined, with the highest level in gall bladder and the lowest level in brain. Triclosan was readily metabolized to triclosan sulfate and glucuronide conjugates, 2,4-dichlorophenol, and hydroxytriclosan. The metabolite profile was tissue-dependent and the predominant route of excretion was the feces (65 - 78%). There were no differences in bioavailability between males and females. Slightly lower absorption was observed with Elizabeth’s Collall, suggesting some oral ingestion due to the mouse’s behavior grooming. (Supported by Interagency Agreement between NCTR/FDA IAG #224-12-0003 and NIH AES12013.)

Confocal Raman Microscopy—The Future for Dermal Absorption Studies.

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The activity and absorption of pesticides into the skin of the user is increasingly becoming a major problem within the Agrochemical industry. Fewer formulations are passing the regulatory process of pesticide approval as a result of unknown compound dermal absorption and toxicity. Currently, 2 gold-standard methods are used to monitor dermal absorption and determine compound flux (μg/cm²/h) across the skin but limitations in measuring dermal absorption most often result in compound failure. Confocal Raman Microscopy (CRM) is novel label free, rapid, optical method which allows compounds to be identified within dermal tissue at depths of ~300 μm. The aims of the investigations were to develop and establish CRM methods for in vitro dermal absorption investigations across ex vivo pig skin. A Lab RA HR CRM was employed for the analysis of a range of concentrations (0.0 - 50.0 mg/mL) of benzoic acid, caffeine (both in 50% (v/v) ethanol (EtOH)) set at the following; IR laser at 785 nm a filter intensity of 100%, a hole 300 μm with a x50 objective. Depth profiling was also performed on fresh pig skin samples from the surface to a maximum depth of ~300 μm (~50 μm increments). CRM spectra for the pig skin investigations demonstrated good reproducibility with some spectral differences noticed. Intensity of the spectra decreased with increased depth as expected. It was observed that a concentration of 50% (v/v) EtOH appeared to mask the signal of prominent identifiable peaks for benzoic acid and caffeine solutions at and below a concentration of 6.25 mg/mL. Compound concentrations equal to or higher than 20.0 mg/mL within a 50% (v/v) EtOH solution could be detected. This data suggests that this masking effect by EtOH requires further exploration to better understand the limitations of the technique. The CRM method has the potential to qualitatively determine dermal compound accumulation within mixed media and solutions ultimately, providing a powerful tool in identifying toxic compounds early on.

Development of an Alternative Model for Assessing Barrier Function and Permeability for Infant Skin.

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Recent published literature of non-invasive measures of skin barrier function of full-term human infants indicates a dynamic maturation process over the first year of life indicative of adaptation from in utero to the external environment. During this period, infant skin may be more vulnerable to penetration of topically applied ingredients, however, the magnitude of differences in non-invasive measures may not correlate to the magnitude of the difference in penetration of topical agents. While thriving demonstrates that a functional skin barrier exists at birth, differences in dermal permeability compared to adult skin could have an impact on the exposure assessment and safety of topically applied ingredients. The purpose of this study was to compare various indicators of skin barrier function (electrical impedance, TEWL), and permeability (tritiated water, 14C-octanol, and 24-h aqueous caffeine) across skin from donors of different ages. Fresh neonatal and pubertal porcine skins (animals culled for other purposes) were collected from a breeding colony. Cryopreserved ex vivo adult human skins were purchased from a skin bank. All measures of skin barrier function and permeability were equally applied to all skin samples. The barrier functions in the porcine skin varied, depending upon the measure. Average impedance indicated parity across age. TEWL and tritiated water flux was lower through neonatal skin versus either mature species. Neonatal skin octanol flux showed parity versus pubescent porcine skin while immaturity versus human cadaver skin. Caffeine absorption ranged from 1.5x to 4.5x higher through neonate skin than seen in either mature species. These data demonstrate that non-invasive measures of barrier function may be indicative, but not definitive, for determining the absorption of topical compounds. The current 10 species safety factor used in risk assessment adequately captures the difference in neonatal barrier function.

Modelling Diffusion of Therapeutic Drugs through the Skin

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A mathematical model accurately simulating diffusion of chemicals through the skin would be useful for predicting percutaneous absorption and aid understanding penetration mechanisms. Models of varying complexity have been validated using
in vivo and in vitro data. In the present study data from the penetration of four drugs was used to test the output of a simple one compartment model based on Fick's 1st law of diffusion.

Methods: Drugs, as off the shelf formulations, were applied to pig skin (500 μm dermatomed slice) in Franz type diffusion static cells using stirred ethanol:water (1:1) as a receptor; surface temperature 32°C. Penetration of hydrocorrosione, capsaicin, ciclosporin and diltiazem were measured in the receptor fluid using Ultra Performance Liquid Chromatography. The experimental permeability constant (Kp) was estimated from the maximum flux rate (assumed to be steady state; Jss) and the applied concentration of drug (C0) using the equation: Kp = Jss / C0. A penetration profile was predicted using a model with the skin as a single compartment of a thickness approximating the stratum corneum and using the experimental Kp to calculate flux of drug from vehicle into the barrier layer and from barrier layer into receptor fluid. A best fit (determined by least squares) was then obtained by adjusting the experimental Kp by a factor which gave an estimate of the accuracy of the model in predicting the experimental data. Results: Model predictions needed to be adjusted by factor of between 1.1 and 1.35 to fit the experimental data as judged by a least squares method and assuming a membrane thickness of between 0.025 and 0.04 mm.

Discussion: This work has demonstrated that a very simple model is able to predict quite closely the penetration of some non-volatile compounds applied to the skin in a vehicle. This work is supported by the US Army Medical Research and Materiel Command under contract W81XWH-08-C-0070. © Dsl Crown Copyright 2012

1047 Prediction of Skin Disposition After Topical Application of Drugs—Simcyp Platform As a Tool for Capturing System Information and Safety Assessment.

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There are reports where topical application of drugs, which is generally considered safe, results in health threatening side effects or drug-drug interactions. Simcyp platform separates the system information from those of drug and uses in vitro in vivo extrapolation approach to predict disposition of the topically administered drugs (Polak 2012). Administration of a single 0.4 mg of erythromycin (topical - 2cm² area of the forearm) with 40 mg simvastatin (oral) is used as a model scenario to assess anticipated interactions mimicking corresponding clinical study (van Hoogdalem 1996). Lotion formulation was simulated by modifying permeability/partition parameters (factor of 50 for stratum corneum and viable epidermis). Simulated residual amounts of erythromycin collected from the skin surface at various time and fraction of the dose recovered from stratum corneum after 6 hours were compared with the observed values. The results were consistent with observations and indicate significant systemic absorption. Subsequently oral simvastatin (40 mg SID) and topical ketocazole (0.4% BID for two weeks, applied on upper arm) were simulated in 100 healthy individuals and by modification permeability/partition parameters (factor of 5 for stratum corneum) to account for occlusion. The average AUC/IAUC and Cmax/Cmax ratios were 1.17 and 1.11 respectively, suggesting safe concomitant use. The maximum simulated ratios were 1.51 (AUC) and 1.52 (Cmax) indicating potential risk of significant interaction in some individuals. This is in agreement with the clinical observations suggesting general safety with rare occurrence of major side effects (e.g. Lareb Q 2009 report; myalgia).


1048 In Vitro Human Epidermal Penetration of IMX-101 Munitions Chemicals.


Dermal exposure to ammunition chemicals is a major concern of the Army in assessing risk during manufacturing and field operations. The US Army Environmental Quality Technology (EQT) program is creating new, insensitive ordnance formulations to replace highly energetic compounds such as RDX and TNT. There are limited data on dermal absorption of munitions chemicals to support health risk assessment. We studied in vitro dermal penetration of a newly-developed munition, IMX-101 (Insensitive Munitions eXplosive) (as a mixture) and its individual chemicals, 2,4-dinitroanisole (DNAN), 3-nitro-1,2,4-triazol-5-one (NQ) and 1-phenyl-1-hexanol (RQ). Human epidermal membranes were prepared from frozen cadaver skin according to OECD guidelines and were mounted in Franz static diffusion cells so that the visceral side was in contact with the receptor fluid. Neat test chemicals at infinite dose (100 mg powder) were carefully placed on the mounted skin in the donor chamber and at different times (1, 2, 4, 6, and 8 hrs) 10 μL of receptor fluid was collected and quantified by HPLC. The rate of diffusion per cm² was calculated for each chemical. The analysis of absorbed chemical in the receptor fluid showed that steady state fluxes of neat NTO, DNAN and NQ were 332, 1.10 and 31.25 μg/cm²/hr respectively. When complete IMX-101 was tested as 100 mg of mixture, the fluxes for NTO, DNAN and NQ independently were 135.9, 1.80, and 236 μg/cm²/2 hr, respectively; NTO and NQ showed about 0.4 and 7 times greater rate of penetration in the combination mix as than individual compounds. These estimated values will be used to determine occupational exposure levels with dermal exposure.

1049 In Vitro Human Skin Penetration of Acetyl Hexapeptide-8 from a Cosmetic Formulation.


Peptides are being incorporated into cosmetic creams marketed as anti-aging/anti-wrinkle products. Acetyl hexapeptide-8 may penetrate into the deep layers of the skin and potentially stimulate biological activity by interfering with neuromuscular signaling as its anti-wrinkle effect. The skin penetration of commercially available acetyl hexapeptide-8 (Ac-EEMQRR-amide) from a cosmetic formulation was determined in human cadaver skin assembled into in vitro diffusion cells. An oil-in-water emulsion containing 10% Ac-EEMQRR-amide was applied to skin at a dose of 2 mg/cm². After a 24 hour exposure, the skin surface was washed to remove unabsorbed peptide. Skin discs were then stripped to determine the amount of peptide in the stratum corneum. Removal of the stratum corneum layers was verified by confocal microscopy. The epidermis was heat separated from the dermis and each skin fraction was homogenized. Skin penetration of Ac-EEMQRR-amide was measured in skin layers by hydrophilic interaction liquid chromatography with tandem mass spectrometry (HILIC-MS/MS) using electron orator fragmentation (ESI) in the positive mode. Stable isotopically labeled hexapeptides were used as internal standards for the quantitation of native hexapeptides to correct for matrix effects associated with ESI. Results (% of applied dose) found the majority of the Ac-EEMQRR-amide was not absorbed and removed by washing (87%). Stratum corneum peptide levels decreased as each layer was removed by tape stripping. Total Ac-EEMQRR-amide found in the stratum corneum was 0.22 % and 0.01% in the epidermis. No peptide was detected in the dermis or receptor fluid. Hairless guinea pig skin penetration of Ac-EEMQRR-amide showed similar results where 0.51% remained in the stratum corneum and 0.01% penetrated into the viable epidermis. This skin penetration data will be useful for evaluating the safety of cosmetic products containing short-chain peptides.

1050 Nevirapine-Induced Skin Rash Is Caused by a Reactive Sulfate Metabolite Formed in the Skin.

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Rationale: Nevirapine (NVP) treatment is associated with significant idiosyncratic immune-mediated skin rash and hepatotoxicity in humans. NVP causes a very similar rash in female Brown Norway rats, and we had previously shown that 12-hydroxylation of NVP is required to induce the rash. In this study we studied the further metabolism and covalent binding of NVP in the rat model and in human skin. Immune activation via IL-1β in the skin was also examined. Methods: An anti-NVP antibody was produced and used in immunohistochemistry studies to detect covalent binding of NVP, 12-OH-NVP, or NVP-sulfate to skin and liver proteins from rodents and humans. In vitro incubations of NVP or metabolites with hepatic microsomes or skin cytosolic fractions from humans, mice, or rats were also performed. The ability of SULT 1A1 to metabolize NVP was studied and cutaneous IL-1β levels were examined by ELISA. Results: Covalent binding was observed in the epidermis of NVP or 12-OH-NVP-treated rats. Major modified bands appeared between 40K-60K. Depletion of PAPs decreased blood levels of NVP-sulfate but did not prevent rash or covalent binding in skin. Topical administration of 1-phenyl-1-hexanol (sulfotransferase inhibitor) prevented rash and covalent binding where applied, and also prevented covalent binding of 12-OH to cytosolic skin fractions and SULT 1A1. IL-1β levels were significantly up-regulated in skin of rats with a rash as well as skin isolates. No skin rash or covalent binding was observed in the skin of NVP treated mice. Conclusions: In contrast to covalent
binding in the liver, which involves direct oxidation to a quinone methide, the re-
active metabolite that covalently binds in the skin is a sulfate. The test substance respons-
able for the rash is formed in the skin of rats and in human skin incubations but not in mice which develop no rash. Further work is being done to confirm the role of IL-1β in NVP-induced skin rash. Funding: Canadian Institutes of Health Research.

1051 In Vitro Discrimination of Skin Sensitizing Haptens and Prohaptens in a Modified KeratinoSens Assay with an Added Metabolic Activation Step.

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Prohaptens are chemicals which may cause skin sensitization after being converted into electrophilic molecules by skin enzymes. In vitro sensitization assays ideally should detect the potential of molecules to act as prohaptens. The metabolic activation system most commonly used in in vitro toxicology is Aroclor-induced rat liver S9 fraction. Even if this system contains higher enzyme activities as compared to those reported in skin, it may serve as a surrogate system to study the potential of chemicals to form reactive, skin sensitizing metabolites. To test this concept, the luciferase induction in KeratinoSens reporter cells treated with chemicals in presence and absence of S9 fractions was measured. Suspected prohaptens such as methylsoueugenol, eugenol, trans-anethol or benz[a]pyrene gave no, or weak, gene induction in absence of S9 fractions, and a strongly enhanced luciferase induction in presence of S9 proving their prohapten status. Haptens like DNCB or cinnamic aldehyde gave a reduced response in presence of S9. We then evaluated whether this metabolic activation assay might be implemented in a tiered screening strategy to screen negatives in the classical KeratinoSens™ assay to enhance sensitivity. To this aim all chemicals classified negative in the classical KeratinoSens™ assay were retested with this activation step. Among the 77 chemicals found as correct-negatives, 74 were also negative in presence of metabolic activation, thus this counter-screen does only slightly reduce specificity. However, based on this comprehensive screening, we found that only a small fraction of the known skin sensitizers need activation by the S9/P450 system, and thus the KeratinoSens™-S9 assay may be useful for the in vitro evaluation of specific classes of potential prohaptens, rather than as a general screening approach. These results will be presented along with results on the predictivity and reproducibility of the KeratinoSens™ assay as accumulated during the prevailisation studies conducted for ECVAM.

1052 Toxicogenomic Characterization of Sensitizer and False-Positive Responses in the Local Lymph Node Assay (LLNA).

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Recent publications have highlighted chemistries which yield false positive re-
ponses in the LLNA when compared with guinea pig and human data. A toxicogenomic approach was applied to provide insight into the molecular and cellular mechanisms that may explain these differential responses. Auricular lymph node gene expression responses were evaluated in female CBA mice exposed to equipotent doses of 9 chemical sensitizers and 7 false positives per the standard LLNA dos-
ing regimen. Lymph nodes were analyzed for HSTJR incorporation on day 6 and gene expression responses on study days 4 and 6. Statistical analyses identified 77 and 473 differentially expressed genes (DEGs) between sensitizers and false posi-
tives on days 4 and 6. Class-based comparison of DEGs showed that the most en-
riched functional categories in the sensitizer-specific subsets were consistent with mechanisms involved in the acquisition of antigen-mediated skin sensitization. Key immune responses at the 4 day time point were restricted to genes involved in early T-cell development including pathways involved in IL2 regulation (IL2 and Egfr), Tbeta and the pre T-cell receptor alpha (Peca). Day 6 responses were more consis-
tent with a mature T-cell response and included genes involved in the DCT/T-cell maturation process such as Il21, Lg3 and Fxyd4. In contrast, false positives exhib-
ited a strong pro-inflammatory expression profile including markers for activated macrophages and neutrophils such as Cds1, Il12b, Mpo, Defa4 and class 1 stefins. Expression of these genes in the absence of dermal irritation suggested these re-
ponses were not solely driven by skin irritation. These gene expression profiles sug-
gest a differential cellular recruitment to the lymph nodes following skin exposure to true sensitizers and false positives and provide a potential new endpoint that could be applied to address false positives and enhance the predictive value of the LLNA.

1053 Inductive Effects on Reactivity of the Contact Allergen Benzoquinone and Its Derivatives to Proteins.

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Benzoquinone (BQ) and substituted benzoquinones (SB) are used for dye and cos-
metics production. BQ is an electrophile known to covalently modify proteins via Michael Addition (MA) but the reactivity, reaction mechanistic domains and aller-
genicity of SB are unknown. Electron withdrawing and electron donating sub-
stituents on BQ were assessed for effects on BQ reactivity and allergenicity. Alternative potential protein binding mechanisms were explored. BQ binding to Cy34 on human serum albumin (HSA) was studied and for BQ and SB reactivity studies, nitrobenzenethiol (NBT) was used as a model nucleophile. Hammert and Taft (HT) constants were used to evaluate the influence of these substituents on chemical reactivity. Both NBT binding studies and HT values demonstrated chlo-
rine SB to be more reactive than methyl and t-buty1 SB. Production of semiquinone radicals from SB and characterization of SB-NBT adducts demonstrated that hap-
tenation may also occur via free radical mechanism which is pH dependent, and vinylc substitution mechanisms, in addition to the predominant MA. BQ and SB dernal allergenicity as evaluated in the murine local lymph node assay (LLNA) was consistent with that predicted by reactivity and HT data. These results demonstrate the effect of substituents on BQ reactivity and allergenicity while suggesting poten-
tial utility of chemical reactivity data and HT values for electrophilic allergen iden-
tification and potency ranking.

1054 Interaction of Para-Phenylenediamine with Human N-Acetylatedtransferases.

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The contact allergen para-phenylenediamine (PPD) is known as a good substrate for N-acetylatedtransferase 1 (NAT1) but we also found that concentrations above 50 μM are accompanied by inhibition of NAT1 activity in human keratinocytes. Here, we investigated the substrate and inhibition characteristics of PPD on NAT enzymes. First we measured whether next to PPD and mono-acetylated PPD (MAPPD) the PPD oxidation product Bandrowski Base (BB) can also be acety-
lated. We therefore incubated PPD, MAPPD and BB with human recombinant NAT1 and NAT2 and found them to be good substrates for both enzymes. NAT1 inhibition characteristic of PPD was further studied using the THP-1 cell line which served as model for antigen-presenting cells. Both PPD and MAPPD are N-acetylated by THP-1 and the acetylation is accompanied by NAT1 inhibition. Concentrations above 1 μM PPD clearly reduced enzyme activity already after 5h while 47% reduction was measured after 24h (200 μM). Independent of the sub-
strate-based enzyme inhihitions, certain compounds are known to oxidize the cat-
alytic cysteine or form adducts with NAT protein. Therefore we studied whether PPD, MAPPD and/or oxidized PPD including BB also interact with recombinant
NAT protein itself in the absence of acetyl coenzyme A. All but MAPPD interact with the protein after 2h and the greatest inhibition was found for oxidized PPD (up to 50%). From these results we can conclude that the observed NAT inhibition may be caused by both substrate dependent and independent effects. NAT1 activ-
ity in PPD-treated THP-1 cells was completely restored after incubation in fresh culture medium for 24h, whereas inhibition caused by 24h treatment with MAPPD could be restored for only 10%.

In sum our data indicate that PPD and its oxidation products can inhibit NAT in
both cases. A complete restoration of NAT activity was only achieved for MAPPD.

1055 Pharmacodynamic Profiling of EGFR Inhibitors in HaCaT Cells.


Skin Rash is a serious adverse effect of EGFR inhibitors observed during anticancer therapy in the clinic and appears to be linked to inhibition of the target pathway. The EGFR inhibitors erlotinib and afatinib were investigated at increasing concen-
trations (0, 0.001, 0.01, 0.1 and 10 μM) in the human keratinocyte cell line
Idiosyncratic drug-induced liver injury (IDILI) typically occurs in a small fraction of patients and often results in removal of otherwise efficacious drugs from the pharmaceutical market. The mechanisms of IDILI are unknown, and animal models of IDILI are few. Several animal models have been developed that suggest that inflammation plays an important role in IDILI. Moreover, the inflammatory mediators tumor necrosis factor alpha (TNFα), interferon gamma (IFNγ), and neutrophil-derived elastase (EL) are essential to the pathogenesis of liver injury in these models. The goal of the present study was to determine if two non-steroidal anti-inflammatory drugs associated with IDILI in humans synergize with TNFα, IFNγ or EL in producing hepatoxicity in vitro. HepG2 cells were treated with either dicyclofenac or sulindac sulfide (the active metabolite of sulindac) and with TNFα, IFNγ or EL or EL + TNFα. Cytotoxicity was significantly increased when cells were treated with dicyclofenac or sulindac in the presence of TNFα compared to cells treated with drug or TNFα alone. IFNγ potentiated the cytotoxicity caused by either drug in the presence of TNFα. EL increased the cytotoxicity caused by dicyclofenac in the presence of TNFα. These results indicate that some of the events that occur in drug-inflammatory stress models in vivo can be reproduced in cultured cells and suggest the possibility of developing in vitro systems that can predict IDILI liability of drug candidates. (Supported by NIH grant RO1DK061315 and T32 GM092715-01A1).

### 1056 Cytokine-Induced Liver Hepatotoxicity of Trovafloxacin in Co-Culture of Hepatocytes and Kupffer Cells.

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Immune-mediated chemical-induced hepatotoxicity, i.e. indirect hepatocellular toxicity resulting from immune cells activating liver inflammatory responses, is often overlooked as a potential mode of action due to unavailability of appropriate in vitro models. Kupffer cells are the largest population of resident macrophages in the liver and thus play a critical role in immune-mediated hepatotoxicity and liver injury. For this reason, we have established a co-culture system of rat primary hepatocytes and Kupffer cells that can be used to model chemical-induced immune responses resulting in acute hepatotoxicity. Hepatocytes and Kupffer cells were cultured for 24-72 hours with various amounts of lipopolysaccharide (LPS) and trovafloxacin (TVX), a compound associated with immune-response-mediated hepatotoxicity in vivo. IL-6 and TNFα were assessed by ELISA or Luminex® bead assays and CYP3A was measured using Luciferin-IPA as a probe substrate to monitor endotoxin-induced functional changes in metabolic capacity. LPS-treated co-cultures showed marked down-regulation of CYP3A activity, which correlated with up-regulation of IL-6 production. This response was blunted in co-cultures treated with LPS and TVX, showing no changes to CYP3A activity and significantly lower production of IL-6. However, TNFα production was unaffected in TVX/LPS-treated co-cultures and markedly up-regulated in Kupffer cell monocultures. LPS/TXV-treated co-cultures also exhibited increased hepatocellular injury, represented by decreased ATP content and increased LDH leakage. This data suggests that shifts in cytokine profiles, especially IL6/TNFα ratios may play a role in immune-mediated chemical-induced hepatotoxicity. Co-culture of hepatocytes and Kupffer cells may represent a powerful in vitro tool to predict adverse liver effects resulting from indirect adaptive immune reactions during chemical exposure. This project was supported by funding from the Long-Range Research Initiative (LRI) of the American Chemistry Council (ACC).

### 1057 Drugs Associated with Idiosyncratic Drug-Induced Liver Injury Synergize with Inflammatory Mediators to Produce Cytotoxicity in a Human Hepatoma Cell Line.

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Idiosyncratic drug-induced liver injury (IDILI) typically occurs in a small fraction of patients and often results in removal of otherwise efficacious drugs from the pharmaceutical market. The mechanisms of IDILI are unknown, and animal models of IDILI are few. Several animal models have been developed that suggest that inflammation plays an important role in IDILI. Moreover, the inflammatory mediators tumor necrosis factor alpha (TNFα), interferon gamma (IFNγ), and neutrophil-derived elastase (EL) are essential to the pathogenesis of liver injury in these models. The goal of the present study was to determine if two non-steroidal anti-inflammatory drugs associated with IDILI in humans synergize with TNFα, IFNγ or EL in producing hepatoxicity in vitro. HepG2 cells were treated with either dicyclofenac or sulindac sulfide (the active metabolite of sulindac) and with TNFα, IFNγ or EL or EL + TNFα. Cytotoxicity was significantly increased when cells were treated with dicyclofenac or sulindac in the presence of TNFα compared to cells treated with drug or TNFα alone. IFNγ potentiated the cytotoxicity caused by either drug in the presence of TNFα. EL increased the cytotoxicity caused by dicyclofenac in the presence of TNFα. These results indicate that some of the events that occur in drug-inflammatory stress models in vivo can be reproduced in cultured cells and suggest the possibility of developing in vitro systems that can predict IDILI liability of drug candidates. (Supported by NIH grant RO1DK061315 and T32 GM092715-01A1).

### 1058 Quantitative Relationship Between Intracellular Lithocholic Acid (LCA) and Toxicity in Rat Sandwich-Cultured Hepatocytes (SCH): Incorporation into a Mechanistic Model of Drug-Induced Liver Injury (DILI).

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One proposed mechanism of DILI is inhibition of the bile salt export pump resulting in cellular accumulation of toxic bile acids (BAs). The purpose of this study was to establish the quantitative relationship between intracellular LCA concentrations and toxicity. This information is essential to link BA kinetics to toxicity in a BA transport inhibition model constructed within DLILysim1,2, a mechanistic model of DILI. Day1 rat SCH in 24-wall well plate were incubated with LCA (25–200μM) for 6, 12, and 24h. LCA toxicity was assessed by LDH release from damaged cells and intracellular LCA levels. Intracellular LCA/metabolites were measured by LC/MS/MS analysis of lysate after hepatocytes were incubated for 5min with Ca+2-free HBSS buffer to open tight junctions using B-CLEAR® technology. LCA, and its taurine (TLCA) and glycine (GLCA) conjugates, accumulated extensively in rat SCH. Cellular LCA concentrations increased with increasing LCA medium concentrations; cellular TLCA and GLCA decreased at LCA doses above 100μM, potentially due to saturation of amicidase. LCA, values estimated based on medium and intracellular LCA concentrations at 12hr were 143 and 1098μM, respectively. An Emax-type relationship was observed between cellular LCA exposure (AUCLCA) and toxicity; half maximal toxicity was observed at AUCLCA of 10.4 and 16.3mMh for the 12 and 24h incubations, respectively. Decreased intracellular ATP concentrations preceded the toxicity measured by LDH. AUCLCA (12-24h) correlated with toxicity at 24 hr (r2=0.97) and AUCLCA (24-48h, r2=0.95). These results provide mechanistic insight that increased cellular exposure of unconjugated LCA leads to ATP depletion and subsequent toxicity. Supported by NIH R01 GM41935 and DILI-sim initiative.

### 1059 Transformation of CO2 Evolution Assays into Multwell Plate Format Assays to Assess Effects of Compounds on Substrate Oxidation in Rat Hepatoma H4IIE Cells.


Perturbations of metabolic processes are involved in several pathologies including cardiac hypertrophy, hepatic vacuolation, and steatosis. Different substrates can be used to distinguish between electron transport chain, the citric acid cycle, β-oxidation,
tion, glycolysis, and amino acid metabolism pathway inhibition. The conventional assay for measuring substrate oxidation is through the use of 14C-labeled substrates and measuring the formation of 14CO2. Historically, these assays are run with a standard 96-well plate for cell incubation with substrate. A nonpeptidic gatekeeper is used to seal the environment to ensure CO2 is collected on a 96-well capture filter plate aligned on top. 14C-labeled glucose, pyruvate, glutamine, butyrate, octanoate, or palmitate were used as substrates and their oxidation was assessed in H4IE cells. Etopoxor, an inhibitor of carboxy-palmitoyl transferase-2, demonstrated a concentration-dependent decrease in palmitate oxidation, however, glucose oxidation increased after treatment suggesting a fuel switching to compensate for the defect in fatty acid oxidation. Methylenecyclopropyl acetic acid (MCPA), a reported inhibitor of medium-chain acyl-CoA dehydrogenase, inhibited both butyrate and octanoate oxidation and had minimal effects on palmitate oxidation. An increase in glucose oxidation was observed after MCPA treatment, similar to that observed with etopoxor suggesting a similar fuel switching. Antiymycin A, an electron transport chain inhibitor, decreased the oxidation of multiple substrates with similar potencies, suggesting inhibition at a point in the metabolic pathway common to all substrates. In conclusion, by transforming the classical CO2 evolution assay into a 96-well format, the effect of various compounds on various substrates can be assessed in an easier, higher throughput manner.

1063 In Vitro Assessment of Drug Induced Liver Injury (DILI) Using a High Content Cellular Imaging System.

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Drug-induced liver injury (DILI) is a leading cause of drugs failing during clinical trials and being withdrawn from the market. In vivo safety testing in pre-clinical species ensures that drugs which enter clinical trials do not cause reproducible and dose-dependent liver injury in man, but is of limited value for exploration of underlying mechanisms and does not assess potential to cause rare idiosyncratic DILI. Implementation of an in vitro cell-based predictive assay early in the drug discovery process would help improve early compound attrition and develop safer drug candidates. We tested compound-treated human hepatocellular carcinoma Hep G2 cells, human induced pluripotent stem cell (iPSC)-derived hepatocytes (iCell® Hepatocytes) and primary human hepatocytes using the Thermo Scientific ToxInsight® IVT platform and DILI Assay Cartridge to determine the hepatotoxicity risk of a compound through the measurement of multiple toxicity biomarkers in individual cells. The compounds we investigated include a number of known hepatotoxic compounds (Ticlopidine, Troglitazone, Nadolixic acid, Mefenamic acid, Phenylbutazone and Aflatoxin B1) and non-hepatotoxic compounds (Aspirin, Fluoxetine and Melatonin). Each compound was tested at eight concentrations in triplicate. The DILI Assay Cartridge allows for the high sensitivity and specificity for predicting hepatotoxicity by simultaneously detecting five multiplexed cellular targets and properties associated with cell loss, cellular redox stress, and mitochondrial stress. The hepatotoxicity prediction using the multiparametric data generated for the test compounds demonstrates high specificity across the three hepatocyte models but varying sensitivity for each hepatocyte model system.
Nonalcoholic fatty liver disease (NAFLD) is the most common form of chronic liver disease in Western society representing a spectrum of liver damage ranging from simple steatosis to the more advanced nonalcoholic steatohepatitis (NASH). Experimental models of NASH include diet-induced models such as the methionine and choline deficient (MCD) diet and genetically obese models such as the leptin-deficient, ob/ob mice. Previous studies have demonstrated alterations in the expression and activity of multiple drug transporters in human NASH, but little information is known regarding the most appropriate animal models for predictive toxicology to extrapolate in vivo hepatic and renal drug disposition data to human NASH populations. The purpose of the current study was to determine which experimental NASH model best recapitulates human pathology as well as alterations in drug transporter expression. Both rat and mouse NASH models were utilized in this investigation and include: MCD diet, atherogenic diet, ob/ob and db/db mice, and fa/fa rats. Histologic and pathologic evaluations confirmed that the MCD and ob/ob androgenic rats, MCD, ob/db and db/db mouse all developed NASH. In contrast, the fa/fa rats did not develop pathological NASH. Hepatic mRNA levels of drug transporters were determined in MCD diet model and the ob/ob and db/db mice best recapitulate the expression profile seen in human NASH. In general, the majority of efflux transporters are induced whereas uptake transporters are decreased. Similar to the liver, efflux transporter expression in the kidney is also induced at both the mRNA and protein level. However, renal uptake transporter expression across these models is differentially altered and is transporter and model-specific. These results suggest that the MCD rat, and the ob/ob and db/db mouse models may be more useful preclinical models to determine the effect of NASH on drug disposition as well as determining extra-hepatic alterations in drug transporter expression.

Regulation of ABC2 Internalization in Nonalcoholic Steatohepatitis.

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Non-alcoholic fatty liver disease affects 30–40% of the United States population and can progress to non-alcoholic steatohepatitis (NASH) which is characterized by an accumulation of fat, oxidative stress, fibrosis and inflammation. It has been previously shown that NASH can have a significant effect on the disposition of many drugs and potentially contribute to the occurrence of adverse drug reactions. NASH-associated altered drug disposition has, in part, been linked to stress-induced alterations of the expression and function of hepatic transporters such as ABC2. ABC2 is located on the canalicular membrane and transports xenobiotic substrates into the bile for elimination from the body. By mechanisms that are not entirely understood, ABC2 has been shown to be internalized in rodent and human NASH livers, as well as the in vitro conditions of acute oxidative stress and lipopolysaccharide exposure. Previous reports have suggested that the activation and cellular localization of radixin, protein phosphatase 1, phosphoinositide 3-kinase, protein kinase A, and protein kinase C (PKC) may be involved in the internalization of ABC2. The purpose of this study is to identify potential mechanisms responsible for ABC2 internalization in NASH. Sprague-Dawley rats were fed a control methionine and choline supplemented diet or a methionine deficient diet (MCD) to model NASH. It was found that in NASH livers there was an increase in the membrane association of the PKCε isoform compared to control livers, but not PKCδ or PKCζ, implicating PKCε as a potential regulator of ABC2 internalization. Additionally, the cellular localization of radixin was not found to change in NASH livers. Further insight into the mechanisms behind functional perturbations of hepatic drug transporters such as ABC2 has implications for improved understanding and identification of targets for alleviating adverse drug reactions in clinical NASH.

Deep Sequencing of miRNA in Livers of Rats Exposed to Hepatotoxins.

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MicroRNAs (miRNAs) are small non-protein coding RNAs. They act as regulators of gene expression, by inhibiting mRNA translation or promoting mRNA degradation, and are involved in diverse physiological and pathological events. Recently we have shown (Koufaris et al. Toxicol Sci: 128, 532, 2012) that the CAR activator phenobarbital (PB) induces deregulation of the hepatic miRNAome in the livers of rats. Next Generation Sequencing (NGS) analysis of small RNAs is a powerful tool that can be used for miRNA expression profiling, and offers a deeper understanding of molecular toxicology. Here we explore the liver miRNAome associated with CAR activation, and to compare this with the effects of the hepatotoxicant di-methylarsinic acid (DMA), which is known to induce mitochondrial perturbation. Male rats were dosed with PB (50 or 1000 ppm) or DMA (4 or 40 ppm) in the diet for 28 days, and sacrificed after 2, 4, 15, and 28. Histopathological evaluation revealed centrilobular hypertrophy at 15 and 28 days after PB treatment and multifocal inflammatory foci with necrosis at 28 days after DMA treatment. Total RNA
was isolated from frozen samples, and library constructs for NGS were generated (TrueSeq Small RNA kit, Illumina). Bisulfite-converted DNA was used in qPCR to validate expression of mRNA families and potential gene targets. Consistent with our previous report, miR200 family was induced by 28 days of treatment with PB at 1000ppm, but not by 50ppm. There was a dose-dependent reduction in the zeb1 and zeb2 transcription factors, which are known to regulate expression of the miR200 family. The zeb1 and zeb2 transcription factors regulate the epithelial to mesenchymal transition, thus PB-mediated miR200 family induction appears to be a homeostatic response to protect the epithelial nature of hepatocytes. NGS permits a more detailed exploration of the response of the miRNAome to hepatotoxicants such as PB, thereby aiding an understanding of the underlying pathways.

### 1070 High-Content Imaging of Rat Hepatocytes Long-Term Cultures to Predict Specific Drug-Induced Liver Toxicity.

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Primary rat hepatocytes were investigated for liver-specific functional changes after repetitive treatments with model liver toxicants at low concentrations for two weeks. Cells were cultured in the Collagen I/Matrigel™ sandwich configuration. High content imaging (HCI) was performed by using the Cellomics™ Arrayscan®. The liver-specific pathologies cholestasis/hyper-bilirubinemia, steatosis and phospholipidosis were investigated by means of fluorescence dyes. Mrp2-mediated canalicular transport, intra-cellular accumulation of neutral lipids and phospholipids were assessed by carboxy-DCFDA, BODIPY or LipidTOX™ Red, respectively. Unspecific cytotoxicity was measured by ATP contents and LDH releases. Cyclosporin A (CsA), Amiodarone (AMD), Rosiglitazone (R Gomez), Chlorpromazine (CPZ), Troglitazone (TGZ), Meflofin (MET), Fenofibrate (FFB), Ibutrofen (IBU), Acetaminophen (ACT), Valproic Acid (VPA) and EMD335823 (EMD) were evaluated at four different concentrations and at five time points (day 1, 3, 7, 10, 14). The results showed that inhibition of canalicular transport, induction of phospholipidosis and steatosis occurred after multiple treatments. Clinical relevant toxicities such as absence of cytotoxicity in the hepatocytes. CsA-CPZ and TGZ treatment caused time- and concentration-dependent inhibition of Mrp2-mediated transport, while AMD and CPZ induced dose-dependent accumulation of phospholipids. CsA was also associated with intracellular accumulation of lipids, whereas VPA did not induce steatosis. Taken together, the current data suggest that the current testing strategy has a high predictive value for compounds inducing hyper-bilirubinemia/cholestasis and phospholipidosis, but not for drug-induced steatosis. In addition, the present work illustrates that HCI is a sensitive and reproducible tool for the evaluations of specific cellular functions and that HCI can be applied in safety profiling of drug candidates. Study is part of the European FP7 PREDICT-IV consortium (grant no 202222).

### 1071 Gene Expression Profiling Identifies Molecular Mechanisms of Tamoxifen-Induced Hepatotoxicity in Mice.

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Tamoxifen is an anti-estrogen drug widely used for the treatment and prevention of estrogen- and progesterone breast cancer. However, despite the beneficial properties of tamoxifen, its use has been associated with the induction of hepatic steatosis.

The goal of this study was to investigate the underlying molecular mechanisms associated with tamoxifen-induced hepatotoxicity. Female and male WSB/Eij mice were fed a diet containing 420 ppm tamoxifen for 12 weeks. The livers of tamoxifen-fed mice exhibited moderate pathomorphological changes, as evidenced by glycogen depletion, despite having profound genotoxic and non-genotoxic alterations. Specifically, the livers of tamoxifen-fed mice were characterized by extensive sex-independent accumulation of tamoxifen-DNA adducts and marked changes in the gene expression. To identify tamoxifen-responsive genes that were differentially expressed between the control and tamoxifen-exposed mice, a t-test, p < 0.01, coupled with a fold-change cut-off ≥ 1.5 was applied. A total of 1245 and 2100 genes were found to be differentially expressed in the livers of tamoxifen-exposed female and male mice, respectively. Interestingly, 361 of the genes were differentially expressed in both female and male mice treated with tamoxifen. Most of these genes were associated with altered lipid metabolism, inflammation, cell death and proliferation, and development of liver fibrosis. These results demonstrate the importance of gene-expression profiling in identifying early hepatic molecular events associated with the chronic administration of tamoxifen.

### 1072 A 3D Co-Culture Model-Based Assay to Assess Liver Kupffer-Cell Activation and Functionality.

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Adverse drug reactions in the liver are one of the major causes of attrition in drug development with both small and large molecules. Novel therapeutic modalities such as antibodies and other biologicals can be taken up by non-parenchymal cells, such as phagocytic cells in the liver (mainly Kupffer cells) and can limit drug exposure and cause liver toxicity. Therefore, evaluating Kupffer-cell uptake can serve as an indirect screen for liver toxicity. However, due to lack of suitable organotypic in vitro models, Kupffer cell uptake has been difficult to study. Here, the use of primary heterotypic 3D rat liver microspheres for assessment of drug-induced Kupffer cell activation is demonstrated. Morphological characterization of rat liver microspheres demonstrated the presence of ED1 and ED2-positive Kupffer cells within the hepatospheres. Electron-microscopy confirmed coating of Kupffer cells between hepatocytes in the rat liver microspheres.

Lipopolysaccharide (LPS) treatment of the rat liver microspheres resulted in increased uptake of acetylated LDL (ac-LDL), whereas incorporation of ac-LDL was reduced when the phagocytosis inhibitor gadolinium chloride was applied, indicating ac-LDL uptake by Kupffer cells is dependent on phagocytosis. An increase in IL-6 secretion was also observed at 48 and 120 hrs post-treatment with LPS, indicating Kupffer cell activation. Together these data demonstrate the suitability of heterotypic rat liver microspheres for assessing Kupffer cell activation and functionality by external stimuli, and highlight the potential of 3D rat liver models for evaluating liver toxicity.

### 1073 Organic Solute Transporter Ostc-Ostβ As a Potential Therapeutic Target for the Metabolic Syndrome.

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The metabolic syndrome has increased in incidence in recent decades, and as a result has become a major health concern. Environmental contaminants have been linked to this, as common toxicants have been found to increase susceptibility to weight gain and/or insulin resistance, key features of the metabolic syndrome. This has resulted in a need for pharmaceuticals that lower fat accumulation and improve insulin sensitivity, and this study identifies a novel potential therapeutic target, the bile acid transporter Ostc-Ostβ.

Our laboratory has generated an Ostc+ mouse in the C57Bl6 background, a strain susceptible to age-related weight gain. Interestingly, Ostc+ mice have a significant reduction in bile acid pool size.

The major purpose of bile acids is to emulsify dietary lipids in the small intestine and facilitate their absorption, but they have also recently been shown to function as signaling molecules that affect lipid and glucose homeostasis. This study aimed to explore the role of Ostc+ in lipid and glucose homeostasis in Ostc+ mice.

A comparison of whole body and fat pad weights of Ostc+/+ and wild type mice at 5 and 12 months of age revealed no differences in these parameters at 5 months, but at 12 months, Ostc+/+ mice had accumulated less fat and had lower total body weights. Similarly, extraction of liver lipids indicated less accumulation of total lipid and cholesterol in hepatocytes in Ostc+/+ mice.

As a potential therapeutic target, Ostc+ mice excrete more lipids in feces, are resistant to age related weight gain and have improved insulin sensitivity and glucose tolerance, suggesting that inhibitors of Ostc+ may be useful in treating the metabolic syndrome.

### 1074 Predicitivity of Drug-Induced Liver Toxicity Using High-Content Screening in 384-Well Cultures of HepG2 Cells and Primary Human Hepatocytes.


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Sponsor: A. Wolf

Pfizer, Groton, MA.

Drug-induced liver injury (DILI) is a major cause of failed drug development and drug withdrawal. Predictive toxicology screening assays are therefore required for early validation of compounds that may cause DILI. Many of these investiga-
study we used multiplexed high content screening, with automated fluorescence microscopy and image analysis, to assess the in vitro toxicity of 122 drugs associated with DILI in man and 22 drugs not reported to cause DILI. Studies were undertaken in HepG2 cells with and without rat S9 fraction and in cryopreserved primary human hepatocytes. Toxicity was determined at 4, 24 or 48 h. The parameters assessed were: cell proliferation (nuclear dyes); apoptosis (anti-activated caspase 3); cell cycle arrest (anti-activated H3); reactive oxygen species generation (H2DFFDA dye); mitochondrial damage (TMRE dye); phospholipid and neutral lipid accumulation. All cell systems provided highly specific discrimination (>90%) between those that caused DILI and non-hepatotoxic drugs. The highest sensitivity was observed in HepG2 cells without S9, using the apoptosis (29%) and phospholipid (22%) endpoints. Addition of S9 did not increase the sensitivity of any of the parameters studied. In human hepatocytes, the 24 h H2DFFDA assay had a sensitivity of 20%, whereas the other endpoints had sensitivities of <11%. We conclude that toxicity studies in HepG2 cells enabled detection of compounds which may cause DILI in humans with high specificity but low sensitivity and that neither the addition of S9 nor the use of human primary hepatocytes provided improved predictivity.

1075 Studying Drug-Induced Cholestasis with a Novel Cellular Model Coexpressing Major Bile Salt Transporters in the Liver.


Drug induced cholestasis and hepatocellular injury are two major manifestations of drug induced liver disease (DILI). Research has shown such injuries are often attributed to inhibition of bile salt transporters in the liver. NTCP, OATP1B1, BSEP and MRP2 have been identified as the major transporters modulating hepatic clearance of bile salts and their conjugates. Recently, we demonstrated that canalicular excetration and intracellular retention of bile salts can be studied with a polarized MDCK cell model concomitantly expressing three major bile salt transporters in the liver, i.e., OATP1B1, NTCP and BSEP. To evaluate the utility of this model in studying drug-induced cholestasis, we tested 20 drugs and their major metabolites that have been reported to cause cholestasis in the human. More than half of the drugs or their metabolites, including trtaglozone, benzothiazolone and bosentan, significantly inhibited B-A canalicular transport (canalicular excetration) of [3H]Taurocholate at clinically relevant concentrations, suggesting that blocking transporter mediated canalicular excetration of bile salts is a major mechanism of drug induced-cholestasis in vivo. Furthermore, a few drugs, such as Rifampicin, remarkably elevated intracellular concentration of Taurocholate (which has been suggested to lead to hepatocellular damage), suggest that at high concentrations, these drugs are more likely to cause hepatocellular atis than the others. Other agents, such as estradiol-17-beta-glucuronide, tamoxifen and pyrazinamide, did not exhibit significant inhibitory effect on Taurocholate transport under the test conditions, which suggests there may be other mechanisms involved. In summary, our study demonstrates that the novel OATP1B1/NTCP/BSEP triple transporter expression model can be a useful economical tool for early-stage screening and for mechanistic study of compounds with cholestasis liability.

1076 Use of a Multiplexed Endpoint Assay Strategy to Assess Model Bioactivated Compounds in Sandwich-Cultured Hepatocytes.

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Sandwich-cultured rat hepatocytes (SCRH) represent a metabolically competent in vitro model for screening and identifying potentially hepatotoxic compounds that correlates well with in vivo results. In this study, we employ a novel strategy for multiplexing a broad complement of metabolic plate-reader assays using SCRH to measure hepatotoxicity of model bioactivated compounds, such as acetaminophen (APAP). This approach yields a profile of early toxicity indicators that provides greater mechanistic insight from fewer experimental repeats. SCRH in 24- or 48-well plates were exposed to 8 concentrations of APAP in the presence or absence of the CYP inducer, dexamethasone (DEX); CYP inhibitor, 1-aminobenzotriazole (ABT); and the glutathione synthesis inhibitor, L-buthionine sulfoximine (BSO). Toxicity was assessed at 24 and 48 hours for Cyp3a activity (Promega P450 GloTM), ATP content (Promega CellTiter-Glo® Assay), lactate dehydrogenase leakage (Promega CytoTox ONE™ Homogenous Membrane Assay), and glutathione content (Promega Glutathione-Glo™ Assay). Resulting ATP, glutathione, and LDH data showed dose- and time-dependent APAP toxicity which was moderated, especially at the 24 hour time point, by ABT and exacerbated by DEX. CYP3A activity was induced up to 4-fold by DEX and significantly inhibited by ABT (5% of DMSO controls). BSO nearly eliminated intracellular glutathione levels by 48 hours and increased hepatocyte APAP sensitivity. Collectively, these data demonstrate the utility of SCRH as a model to assess bioactivated compounds for hepatotoxicity. The multiparameter assay approach offers greater mechanistic insight into the nature and timing of the toxicity.

1077 Cytotoxic Interaction of Inflammagens with Trovafloxacin in Spheroid Cultures of Rat Liver Cells.

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Idiosyncratic, drug-induced liver injury (IDILI) typically escapes detection in preclinical safety evaluation of drug candidates due to paucity of predictive models. Recently, model have arisen in which liver injury occurs in rodents treated with trovafloxacin (TVX) or other IDILI-associated drugs and coexposed to an inflamagen such as bacterial lipopolysaccharide (LPS). The hepatocellular injury depends on tumor necrosis factor-alpha (TNF) and possibly other cytokines released by nonparenchymal cells (NPCs) and therefore cannot be reproduced in monocultures of hepatic parenchymal cells (HPCs). Accordingly, the cytotoxicity of TVX in the presence and absence of LPS was evaluated in 3-dimensional (3D) spheroid cultures (in Spherosphere) containing equal numbers of HPCs and NPCs and compared to data obtained from HPCs cultured in standard sandwich culture. Spheroids or standard HPC cultures were exposed to various concentrations of TVX (10–100 μM) alone or with LPS (10 μg/ml) or TNF (10 ng/ml) for 24 h. TNF was detected in the culture medium of spheroids cotreated with LPS and TVX, but not with either agent alone, and no TNF was detected in HPC cultures treated similarly. Cytotoxicity, as assessed by decreases in intracellular ATP content, was evident only in spheroids cotreated with LPS and TVX. In contrast, treatment of either spheroid or HPC cultures with TNF in the absence or presence of TVX did not result in consistent cytotoxicity. Levofloxacin (LVX) is in the same pharmacological class as TVX, but has far less IDILI liability; interestingly, there was no cytotoxic interaction with LVX and LPS or TNF. These results show that LPS-stimulated inflammatory mediator production and cytotoxicity occurred in 3D spheroid cultures containing NPCs cotreated with TVX. 2D-sandwich HPC cultures were less sensitive to cytotoxicity from the TVX-inflammation interaction, suggesting that spheroids may afford advantages in evaluating interactions between drugs and inflammatory mediators.

1078 Trovafloxacin Induces Cell Cycle Arrest and Sensitizes HepG2 Cells to TNF-Induced Cytotoxicity by an ATR-Dependent Mechanism.

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Trovafloxacin (TVX) is an antibiotic associated with idiosyncratic hepatotoxicity in humans, though the mechanism for this toxicity is unknown. In mice, TVX is not hepatotoxic by itself but synergizes with tumor necrosis factor-alpha (TNF) to produce liver injury. Moreover, treatment with combinations of TVX and TNF were cytotoxic to HepG2 human hepatoma cells, whereas neither TVX nor TNF alone caused cytotoxicity. It was demonstrated previously that TVX is a weak inhibitor of ataxia telangiectasia DNA-p kinase (ATK), as well as the tumor suppressor protein p53, all play a role in regulating cell cycle. Pharmacologic inhibitors of ATM, DNAPK, and p53 had no significant effect on the cytotoxicity caused by TNF/TNF treatment, whereas an inhibitor of ATR attenuated cytotoxicity. Taken together, these data suggest that interaction of TVX/TNF with ATM/p53/ATR and ATP and DNA kinase (DNAPK), as well as the tumor suppressor protein p53, all play a rule in regulating cell cycle. Pharmacologic inhibitors of ATM, DNAPK, and p53 had no significant effect on the cytotoxicity caused by TNF/TNF treatment, whereas an inhibitor of ATR attenuated cytotoxicity. Taken together, these data suggest that interaction of TVX/TNF with ATM/p53/ATR and DNA kinase (DNAPK), as well as the tumor suppressor protein p53, all play a role in regulating cell cycle. Pharmacologic inhibitors of ATM, DNAPK, and p53 had no significant effect on the cytotoxicity caused by TNF/TNF treatment, whereas an inhibitor of ATR attenuated cytotoxicity. Taken together, these data suggest that interaction of TVX/TNF with ATM/p53/ATR and DNA kinase (DNAPK), as well as the tumor suppressor protein p53, all play a role in regulating cell cycle.
Alcoholic liver disease (ALD) and non-alcoholic fatty liver disease (NAFLD) are histologically similar diseases as both induce fatty liver (hepatostasis) through mechanisms not yet fully understood. MicroRNAs (miRNAs), small non-coding RNAs are involved in regulation of various biological processes and dysregulated microRNA expression is observed in various diseases including fatty liver. To understand the role of miRNAs in inducing hepatostasis, miRNAs were analysed in the livers of both ALD and NAFLD models. Male Fisher-344 rats were fed with or without 5% alcohol via Lieber-DeCarli diet for one, two or three months or L-aminoacid rodent diet with or without methionine-choline for one week. We observed differential expression of eleven miRNAs in both treatment models. Increased expression of miR-122 and miR-34a was observed in ethanol treated rats as time progressed while miR-122 expression decreased and no change in miR-34a expression was observed in rats fed methionine-choline deficient (MCD) diet. Contrary, expression of miR-451 was increased in both ethanol fed rats with time as well as MCD diet fed rats. However, the increase in miR-451 expression was more pronounced in MCD diet fed rats. miR-122 is known to be involved in hepatic lipid metabolism and sterol synthesis, miR-34a is involved in lipid metabolism, cell cycle and apoptosis while miR-451 is involved in regulation of insulin dependent (glucogenonpheny) pathway. Our miRNA expression data also supported the differential regulation of genes involved in lipid metabolism such as sterol regulatory element-binding protein (SREBP), AMP-activated protein kinase (AMPK), peroxisome proliferator-activated receptor alpha (PPAR-α), sirtuin-1 and acyl-CoA synthetase long-chain family member 1 (ACSL-1). In conclusion, our studies show that differential expression of miRNAs results in steatosis in both models but could be via different mechanisms.

HepaRG (HRG) cells are a next generation version of HepaRG (HRG), a human hepatoma cell line with substantial similarity to primary human hepatocytes (HH). We characterized cryopreserved EP from Biopredic International to evaluate their suitability as a HH surrogate for in vitro toxicity and ADME studies, and for comparison to HRG (Biopredic). The DMSO content of the EP cryopreservation medium was low enough for compatibility with a no spin thaw/recovery protocol that yielded about 90% viable cells (HRG required DMSO removal). One cryopreserved vial of EP provided for two full confluent 96 well plates of cells, which exceed the yield from typical commercially sourced vials of HH. EP cell viability was relatively constant over a period of 7 days in culture. Like HRG, EP established an adherent co-culture of hepatocytes and cholangiocytes but EP had a higher hepatocyte/cholangiocyte ratio. Consistent with this hepatocyte enrichment, EP also had higher activity for CYP1A2, 2D6, 2B6 but similar CYP3A4 activity, and these activities were within the range typically observed in HH. The initial EP CYP3A4 activity remained elevated for the first 24 hours, but then declined to a lower yet still robust level that averaged about 25% of the initial activity over the next 6 days. As a marker of metabolic capacity, the sustained CYP3A4 activity suggests EP will provide a good model for acute or extended drug metabolism studies. Omeprazole, phenobarbital and rifampicin respectively caused substantial inductions of CYP1A2, 2B6 and 3A4 activities with a more pronounced CYP1A2 response in EP compared to HRG. In common with HH, dose-dependent aflatoxin-B1 toxicity was observed with HRG and EP, indicating their competence for detecting metabolism-based toxicities. In summary, EP provides a convenient model for hepatoxicity, metabolism and CYP induction studies with the reproducibility inherent to a cell line. For the purposes of this study EP is a viable HH surrogate with enhanced metabolic capacity and convenience over HRG.

Drug-induced liver injury (DILI) is difficult to predict using classical in vitro assays and regulatory animal studies leading to interruption of clinical trials or even to market withdrawal after commercialisation. Consequently, drug-induced mitochondrial dysfunction should be detected early, ideally during pre-clinical screening of potential candidates could be done by a validated disease model. The present study disclosed for the first time the identification of several drugs triggering drug-induced liver injury (DILI) and a usable model for acute or extended drug metabolism studies. Omeprazole, a marker of metabolic capacity, the sustained CYP3A4 activity suggests EP will provide a good model for acute or extended drug metabolism studies. Omeprazole, phenobarbital and rifampicin respectively caused substantial inductions of CYP1A2, 2B6 and 3A4 activities with a more pronounced CYP1A2 response in EP compared to HRG. In common with HH, dose-dependent aflatoxin-B1 toxicity was observed with HRG and EP, indicating their competence for detecting metabolism-based toxicities. In summary, EP provides a convenient model for hepatoxicity, metabolism and CYP induction studies with the reproducibility inherent to a cell line. For the purposes of this study EP is a viable HH surrogate with enhanced metabolic capacity and convenience over HRG.

Activation of Constitutive Androstane Receptor (CAR) has been shown to protect against bile acid (BA)-induced liver injury. However, the effect of CAR activation on bile flow, BA profile, and BA synthesis and transport genes in liver is not well understood. Therefore, the purpose of the present study was to determine the effect of CAR activation on BA homeostasis. The CAR ligand 1,4-Bis-[2-(3,5-dichloropyridyloxy)]benzene (TCPOBOP) was administered intraperitoneally (3 mg/kg). We found the male C57BL/6N rats treated with TCPOBOP for 4 days, followed by collection of serum, liver, and bile. BAs of serum and bile were quantified by UPLC-MS/MS. TCPOBOP induced a CAR-dependent increase in bile flow by 67%. TCPOBOP increased biliary excretion of two of the three quantitatively major bile acids in mice, TUDCA (3.5-fold) and TCDCA (8%); as well as the quantitatively minor bile acids TUDCA (38%) and TCDCA (224%). Additionally, TCPOBOP increased biliary excretion of unconjugated bile acids by 74%. These increases in biliary excretion correlated with increased liver mRNA of the two major canalicual efflux transporters MRP2 (3.2-fold) and Bsep (2.4-fold). Interestingly, TCPOBOP-induced BA synthesis genes Cyp7a1 (4.3-fold) and Cyp27a1 (110%). Also in livers of TCPOBOP-treated mice, the sinusoidal efflux transporters Mrp3 (17-fold) and Mrp4 (14-fold) were markedly increased. Surprisingly, there were no changes in serum concentrations of total BAs, taurine-conjugated BAs, unconjugated BAs, or primary BAs, but the serum secondary BAs were decreased 61%. In summary, while CAR activation has relatively minor effects on serum BA profile, CAR activation increases bile flow and biliary excretion of TUDCA, TCDCA and total unconjugated BAs. (R01ES009649, R01DK081461, F32DK092069)

MicroRNA Regulation of Alcoholic and Nonalcoholic Fatty Liver Disease.

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Failure in regulation of bile acid (BA) homeostasis contributes to several liver diseases. It is generally accepted that hepatic FXR-SHP and intestinal Fgf5/19 regulate BA homeostasis. However, the contribution of other hepatic transcription factors to BA homeostasis is not clear. The adaptive role of the aryl hydrocarbon receptor (AhR) in protecting against BA related liver damage is unknown. The AhR tors to BA homeostasis is not clear. The adaptive role of the aryl hydrocarbon receptor (AhR) in protecting against BA related liver damage is unknown. The AhR tors to BA homeostasis is not clear.

Mitochondria to Predict Drug-Induced Liver Injury in Human.

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Drug-induced liver injury (DILI) is difficult to predict using classical in vitro assays and regulatory animal studies leading to interruption of clinical trials or even to market withdrawal after commercialisation. Consequently, drug-induced mitochondrial dysfunction should be detected early, ideally during pre-clinical screening of potential candidates could be done by a validated disease model. The present study disclosed for the first time the identification of several drugs triggering drug-induced liver injury (DILI) and a usable model for acute or extended drug metabolism studies. Omeprazole, a marker of metabolic capacity, the sustained CYP3A4 activity suggests EP will provide a good model for acute or extended drug metabolism studies. Omeprazole, phenobarbital and rifampicin respectively caused substantial inductions of CYP1A2, 2B6 and 3A4 activities with a more pronounced CYP1A2 response in EP compared to HRG. In common with HH, dose-dependent aflatoxin-B1 toxicity was observed with HRG and EP, indicating their competence for detecting metabolism-based toxicities. In summary, EP provides a convenient model for hepatoxicity, metabolism and CYP induction studies with the reproducibility inherent to a cell line. For the purposes of this study EP is a viable HH surrogate with enhanced metabolic capacity and convenience over HRG.

CAR Activation Increases Bile Flow, Excretion of Major Bile Acids, and Bile Acid Synthesis, As Well As Transport Genes in Mice.

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Failure in regulation of bile acid (BA) homeostasis contributes to several liver diseases. It is generally accepted that hepatic FXR-SHP and intestinal Fgf5/19 regulate BA homeostasis. However, the contribution of other hepatic transcription factors to BA homeostasis is not clear. The adaptive role of the aryl hydrocarbon receptor (AhR) in protecting against BA related liver damage is unknown. The AhR may be critical for mediating an adaptive response that involves induction of genes responsible for the metabolism or transport of BAs. To study the role of the AhR in BA homeostasis, sera, livers, and bile of 12-week-old male wild-type and AhR-null mice were collected, and BA concentrations in serum and bile, as well as the mRNA expression of hepatic BA synthesis and transport genes were quantified. Surprisingly the lack of AhR increased serum total BAs (62-fold), taurine conjugated-BAs (52-fold), and unconjugated-BAs (113-fold). Similarly the biliary excretion of total BAs (2.3-fold), taurine conjugated-BAs (2.3-fold), and unconjugated-BAs (3.8-fold) increased in AhR-null mice compared with controls. In livers of...
AMPK is a key sensor and regulator of glucose, lipid, and energy metabolism throughout the body. Activation of AMPK improves metabolic abnormalities associated with metabolic diseases including obesity and type-2 diabetes. AMPK phosphorylation inhibits sterol regulatory element-binding protein-1 (SREBP-1) activity which plays an important role in lipid metabolism. In this study, we established in vitro screening methods for potential activators of AMPK from herbal extracts to develop the inhibitor of hepatic steatosis. First, the phosphorylation of AMPK and its substrate ACC-1 were screened by In-cell-western assay which is an immunocytochemical assay performed in 48 well-microplate format and then confirmed them using Western blot in HepG2 cells. Oil Red O staining is performed to observe the accumulation of lipid droplets in cell and then tested the expression of genes related lipid synthesis and metabolism. When we screened over 50 herbal extracts, we found three extracts which were significantly increased the phosphorylation of AMPK and its substrate ACC-1 in a dose-dependent manner. They also reduced the palmitic acid induced lipid droplets accumulation determined by Oil Red O staining. Among them, one extract was selected and observed SREBP-1 transcriptional activity using a SRE-luc reporter gene assay and SREBP-1 target genes expression using RT-PCR. In this study, we could identify the natural extracts for the development of AMPK activator for diabetes and the metabolic syndrome.

Oleanolic acid (OA) is a triterpenoid that exists widely in fruits and medicinal herbs. OA is effective in protecting against hepatotoxins. However, we recently found that whereas a low dose of OA is hepatoprotective, higher doses and long-term use of OA produce cholestasis. This study characterized OA-induced cholestatic liver injury. Adult C57BL/6 mice were given OA at doses of 0, 50, 100, 200, and 300 μmol/kg, sc (suspended in 2% Tween-80 saline) for 5 days, and cholestatic liver injury was observed at doses of 200 μmol/kg and above, as evidenced by increases in serum total bilirubin, serum activities of alanine aminotransferase and alkaline phosphatase, as well as by histopathology with feathering-like degeneration indicative of cholestasis. Serum bile acid (BA) concentrations were dramatically increased not only for unconjugated bile acids (CA, CDCA, DCA, αMCA and βMCA), but also for tauro conjugated bile acids (TCA, TCDCA, TDCA, TUDCA, and T/αMCA), as determined by UPLC-MS/MS. The mRNAs of hepatic bile acid uptake transporter for unconjugated BAs Oatp1b2 and conjugated BAs Ntcp, as well as canicular efflux transporter Bsep were decreased in a dose-dependent manner. OA decreased OSTα but increased OSTβ. Hepatic uptake transporter Oatpl1a1, 1α, and 2b1 were suppressed, while the efflux transporters Abca1, Abt1, and Bcp were increased. OA decreased the mRNA of the nuclear receptors CAR, FXR, PPARα but had no effects on AHR and PXR. The mRNA of the BA biosynthesis limiting enzyme Cyp7a1 was decreased. Taken together, higher doses of OA produces cholestatic liver injury in mice and this effect appears to be associated with the alteration of liver transporters resulting in disruption of BA homeostasis.

The diagnosis of drug-induced liver disease is an intriguing question oftenaccompanied by insufficient clinical data and the difficulty of confirming histopathology after exposure to toxins. This study investigated the histopathological changes induced by several different bioactivation-dependent hepatotoxins, such as, acetaminophen (an analgesic; 500mg/Kg, ip), carbon tetrachloride (CCL4; a hepatotoxin), dimethylnitosamine (DMN; a potent carcinogen), doxorubicin (DOX; an antineoplastic agent), furasemide (FUR; a loop diuretic) and streptozocin (STZ;
an inducer of hyperglycemia) to characterize most common and most uncommon pathological traits in order to hypothesize whether a subset of lesions can be used to diagnose toxicity profile of a particular toxin. Mice were administered CCl4 (1ml/Kg, po), DMN (50 mg/Kg, ip), DOX (60 mg/Kg, ip), Fur (500 mg/Kg, ip) and STZ (100 mg/Kg, ip 2 consecutive doses) and sacrificed 24-72 hours later. Blood was collected for serum chemistry and liver sections for histopathology. PAS or H&E stained liver sections were examined under a brightfield microscope to determine similarities concerning: sinusoidal dilation, centrilobular necrosis, excessive vacuolization, cells with apoptotic morphologies, microvesicular steatosis, ballooning degeneration, pericentral or periportal or indiscriminate necrosis, glycogen depletion trend, inflammation, and formation of ground glass hepatocytes. Results indicated that all the toxins, except for massive liver injury with limited overlapping profiles. The most common features were indiscriminate glycogen depletion and ballooning of hepatocytes. Results also indicated various other characteristic morphological changes such as macrovesicular steatosis, bridging necrosis, portal tract fibrosis and some unique but yet to be reported feature induced by DOX, i.e., prominent sinusoidal dilation. Thus, our study made a serious effort in differentiating and profiling the varied patho-morphological changes induced by diverse hepatotoxins. [Suppo. by Dept. of PScs, AMS Coll of Pharm & HScs]

1089 Cholesterol Liver Effects in Healthy Human Subjects.

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Cholesterol is an orally administered, an ion exchange non-absorbable resin with a high affinity for biliary acids used as a cholesterol lowering agent as well as to accelerate clearance from the body of xenobiotics and to treat itching in patients with advanced cholestatic liver disease. This treatment has been associated with elevations in serum alanine aminotransferase (ALT), a sensitive biomarker of liver injury, but has never been associated with clinically important liver injury. A recent clinical trial in healthy adults receiving cholestyramine provided an opportunity to use current serum biomarkers to investigate mechanisms underlying this phenomenon. After a double blind, placebo controlled study cholestyramine (8 gm/tid X 11 days) was administered to healthy adult subjects to accelerate clearance of the new drug candidate given in the active arm. During cholestyramine therapy, eleven subjects previously treated with placebo experienced elevations in serum ALT levels exceeding three-fold the upper limit of normal (3x ULN; mean 6.9 fold; range 3-28 fold). Serum samples from those subjects were assayed for additional mechanistic biomarkers. Compared to predose baseline values, there was a significant (p<0.05) 13-fold mean increase in mIR-122 confirming a liver source for the serum ALT. Mean serum levels of GLDH (8-fold), cytochrome 18, and HMGBl (1.7 fold) were elevated supporting ongoing necrosis. Caspase-cleaved cytokeratin 18 was also increased (1.7 fold) supporting apoptosis. Serum ALT elevations observed during cholestyramine treatment reflect both hepatocellular necrosis and apoptosis. The wealth of clinical data available with cholestyramine indicate that this effect on the liver is self-limited and not a significant safety concern.

1090 Sterile Inflammation and Neutrophil Activation during Injury Resolution following Acetaminophen Overdose in Mice and Humans.


Following acetaminophen (APAP)-overdose, there is a robust inflammatory response triggered by the release of cellular contents from necrotic hepatocytes into systemic circulation which recruits neutrophils (PMNs) into the liver. PMNs do not contribute to APAP-induced injury in mice, but their role and the role of NADPH oxidase, in injury resolution is unclear. C57BL/6 mice were treated with APAP and were euthanized (6h - 72h) for assessment of liver injury and regeneration. Informed and consented human APAP-overdose patients (ALT >800U/L) had serial blood draws during injury and recovery phases for determination of liver function tests (LFTs) and flow cytometric analysis. Blood PMNs (mouse, human) and liver PMNs (mouse) were evaluated by flow cytometry (CD11b, reactive oxygen generating primers, phagocytosis). To test the importance of phagocyte ROS production gp91phox-/- (NADPH oxidase defective) mice were compared to C57BL/6 mice for hepatocyte proliferation and injury resolution. PMNs in the peripheral blood of mice showed increasing activation status during and after the peak of injury but returned to baseline levels prior to injury resolution. Hepatic sequestered PMNs showed an increased and sustained CD11b expression, but no ROS priming was observed. Confirming that NADPH oxidase is not essential for recovery, gp91phox-/- mice following APAP-overdose displayed no difference in injury resolution. APAP-overdose patients also showed increased PMN activation status after the peak of injury, and PMNs remained in an activated state even as LFTs began to normalize. The time course of PMN activation status in the peripheral blood was similar in mice and humans, and markers of activation were increased and sustained well after active liver injury had subsided. The similar findings between surviving patients and mice indicate PMN activation may be a critical event for injury resolution following APAP-overdose in man, but not a contributing factor to APAP-induced injury.

1091 Ethyl Tertiary -Butyl Ether Induces Oxidative Stress and 8-OHdG Formation in the Liver of F344 Rats via Activation of CAR, PXR and PPAR Nuclear Receptors.

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To elucidate the mode of action (MOA) of hepatotoximogenicity in rats for non-genotoxic chemical ethyl tertiary-butyyl ether (ETBE), male F344 rats were treated with ETBE at doses of 300 and 2000 mg/kg/day by gavage and 500 ppm pphenobarbital (PB) in diet for the comparison analysis for 1 and 2 weeks. Significant increases of hydroxyl radicals and P450 total content in liver and kidney were observed in rats treated with ETBE at doses of 300, 2000 mg/kg/day ETBE and PB groups accompanied the accumulation of P450 isoenzymes CYP2B1/2, CYP3A1/2 and CYP2C6 in the cytoplasm of hepatocytes. Specific up-regulation of CYP2E1 and CYP1A1 was obvious in the high dose ETBE group. Consipicuous elevation of 8-hydroxydeoxyguanosine (8-OHdG) and apoptosis in the liver tissue observed in 2000 mg/kg/day ETBE and PB groups was associated with suppression of Ki-67 positive-cell index, cyclin D1 mRNA expression and down-regulation of 8-OHdG repair enzyme, DNA glycosylase 1 (Ogg1) after 2 weeks of application. Results of QSTAR LC/MS/MS and IPA analyses indicated that upstream regulators of gene expression altered by ETBE altered included CAR, PXR and PPAR. High dose ETBE induced peroxisome proliferation in hepatocytes at week 2 detected by TEM. These results indicated that MOA of ETBE hepatotoxicity in rats is due to induction of oxidative stress and 8-OHdG formation, subsequent cell cycle arrest and apoptosis, suggesting regenerative cell proliferation, predominantly induced by activation of CAR and PXR nuclear receptors by a mechanism similar to that of PB, and furthermore peroxisome proliferation by specific activation of PPAR. The MOA of ETBE hepatotoxicity in rats is indicated to be not relevant to human.

1092 Expression Profiles of Hematopoietic Stem Cell, Endothelial Cell, and Myeloid Cell Antigens in Spontaneous and Chemically Induced Hemangiomias and Hemangioendotheliomas in Mice.

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It is unclear whether the process of spontaneous and chemically induced hemangiomia (HA) and hemangioendothelioma (HS) formation in mice involves the transformation of differentiated endothelial cells (ECs) or recruitment of multipotential bone marrow-derived hematopoietic stem cells or endothelial progenitor cells (EPCs) which show some degree of endothelial differentiation. In the present study, immunohistochecmic staining for hematopoietic stem cell markers (CD45 and CD34), EC markers [VEGFR2, CD31, and Factor VIII-related antigen (FVIII RAg)], and a myeloid lineage marker (CD14) was used to assess the origin of HA and HS in mice and compare with results of canine and human EC tumors reported in previous literature. We showed that the staining for CD45, FVIII RAg, and CD14 was negative in spontaneous and chemically induced HA and HS, whereas cells stained positive for CD34, VEGFR2, and CD31 in these tumors. These results indicate that mouse HA and HS are composed of cells derived from EPCs expressing CD34, VEGFR2, and CD31, but not FVIII RAg. The lack of
CD34 expression suggests that mouse vascular tumors may arise from EPCs that are at a stage later than hematopoietic stem cells (HSCs). Since FVIII RAg expression is known to occur later than CD31 expression in EPCs, our observations may indicate that these tumor cells are arrested at a stage prior to complete differentiation, in contrast to vascular tumor cells in humans and canines, which express both CD31 and FVIII RAg. In addition, unlike human infantile HA or canine HS, myeloid lineage cells do not appear to contribute to HA and HS formation in mice. Our results may indicate that mouse HA and HS may arise by a different mechanism than canine or human EC tumors.

1093 Role of the Circadian Clock in the Differential Susceptibility of Rat Strains to Mammary Carcinogenesis.

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Disruption of circadian rhythm is associated with increased risk of breast cancer incidence and malignancy. DNA damage responsive and repair pathways are controlled by and also modulate the molecular circadian oscillators. In order to investigate the role of the circadian clock on the susceptibility to mammary tumor, we compared circadian expression patterns of genes involved in circadian rhythm and DNA damage responsive and repair pathways in mammary glands of susceptible Fisher 344 (F344) and resistant Copenhagen (COP) rats. The mammary tissue of susceptible F344 rats showed a 4-hr delay in the rhythmic expression of clock genes (e.g., Per2 and Bmal1) and much higher amplitude on Rev-ErbAβ expression compared to resistant COP rats. In mammary glands of COP rats, 45% of DNA responsive and repair genes (total 82 genes) were up-regulated in a circadian pattern with peak expression levels over 2 folds at Zeitgeber Time (ZT)12 (lights off) compared to ZT0 (lights on). By contrast, in the susceptible F344 rats, 30.5% of these genes showed a circadian pattern of down-regulation, with only one gene being up-regulated in the active phase. Exposure to a single carcinogenic dose of N-nitroso-N-methylurea (NMU) disrupted the rhythmic expression of Per2 in mammary gland of F344 rats after two days and further abolished after 30 days. In contrast, NMU induced a significant increase in the rhythmic expression of Per2 gene in the Cop rats in 2 days, and the increase was sustained for at least 30 days post exposure. Moreover, the ratio of NAD+/NADH and NAD+–dependent Sirt1 activity were also significantly increased in NMU-treated COP, but were dramatically decreased in treated F344 rats. Taken together, we speculate that the genetic block to tumor progression in COP is, at least in part, due to the ability to maintain or enhance circadian rhythm, and hence DNA repair upon exposure to carcinogens. (Supported by NIEHS grants, U19ES011387, P30ES007033, and P30ES005022).

1094 Transcriptomic Profiling of Hepatoblastomas and Associated Hepatocellular Carcinomas in B6C3F1 Mice.

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The cell of origin of hepatoblastoma (HB) is unknown; it is thought to be a transformed hepatocyte, an oval cell, or a multipotent hepatic progenitor cell. Many investigators believe that HBs arise solely within hepatocellular carcinoma (HCC), and therefore are a more malignant form of HCC. Comparison of differential gene expression changes in HB to HCC in which they arise may identify alterations specific to each tumor type, as well as alterations shared by both tumor types. Such studies may provide information on the possible cell of origin and pathogenesis of these tumors. In the current study, frozen samples from chemically-induced HB, and HCC were analyzed. Principal component analysis of the normalized transcriptomic data showed that the HB samples clustered distinctly from HCC. When compared to AN, there were increased numbers of differentially expressed genes in HB (10,386 genes) compared to HCC (1,073 genes). Genes involved in Wnt/β-catenin signaling, hepatic embryonic cell-like phenotype and genomic imprinting were significantly altered in HB compared to HCC, pathways important in human HB but not much different from regular insulin. Therefore, these data suggest that insulin glargine is similar in mitogenic potency to AspB10 and IGF1, and detemir. To separate INSR versus IGF1R activation motilities of these analogs and cancer incidence in diabetic patients. Different insulin analogues and cancer incidence in diabetic patients. Different insulin analogues might affect the affinity and activation of these analogues towards the insulin receptor (INSR) and the insulin-like growth factor 1 receptor (IGF1R). A switch towards higher IGF1R affinity is likely to emphasize cellular signalling pathways that promote mitogenesis rather than glucose metabolism. To investigate this hypothesis we have performed in vitro exposure experiments with several insulin analogues including glargine, AspB10, aspart, glulisine, lispro and detemir. To separate INSR versus IGF1R activation motilities of these analogues, we used human breast cancer cell lines (MC7) that either ectopically express the INSR (A or B isoform) in conjunction with a stable knockdown of the IGF1R, or the IGF1R in conjunction with a stable knockdown of the INSR. Using Western blots we measured the activation of INSR and IGF1R as well as downstream mitogenic signalling cascades including ERK and Akt. We revealed differences in activation patterns between the different insulin analogues tested. While insulin primarily acted through the INSR, the mitogenic signalling activation pattern of the long-acting insulin glargine was highly similar to that of known mitogenic compounds like insulin AspB10 or insulin like growth factor 1 (IGF1) and acted mainly via the IGF1R. These findings correlated with the proliferative effect of insulin glargine: at low insulin glargine concentrations a clear increase in cell proliferation was observed. This occurred despite the fact that the EC50 for the proliferative effect in human HB in vitro is not much different from regular insulin. Therefore, these data suggest that insulin glargine is similar in mitogenic potency to AspB10 and IGF1, at least at low and presumably physiological concentrations.

1096 Differential Activation of Mitogenic Signaling through the Insulin Receptor or the Insulin-Like Growth Factor 1 Receptor of Various Clinical Relevant Insulin Analogs.

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Several epidemiological studies suggest an association between the use of some insulin analogues and cancer incidence in diabetic patients. Different insulin analogues might affect the affinity and activation of these analogues towards the insulin receptor (INSR) and the insulin-like growth factor 1 receptor (IGF1R). A switch towards higher IGF1R affinity is likely to emphasize cellular signalling pathways that promote mitogenesis rather than glucose metabolism. To investigate this hypothesis we have performed in vitro exposure experiments with several insulin analogues including glargine, AspB10, aspart, glulisine, lispro and detemir. To separate INSR versus IGF1R activation motilities of these analogues, we used human breast cancer cell lines (MC7) that either ectopically express the INSR (A or B isoform) in conjunction with a stable knockdown of the IGF1R, or the IGF1R in conjunction with a stable knockdown of the INSR. Using Western blots we measured the activation of INSR and IGF1R as well as downstream mitogenic signalling cascades including ERK and Akt. We revealed differences in activation patterns between the different insulin analogues tested. While insulin primarily acted through the INSR, the mitogenic signalling activation pattern of the long-acting insulin glargine was highly similar to that of known mitogenic compounds like insulin AspB10 or insulin like growth factor 1 (IGF1) and acted mainly via the IGF1R. These findings correlated with the proliferative effect of insulin glargine: at low insulin glargine concentrations a clear increase in cell proliferation was observed. This occurred despite the fact that the EC50 for the proliferative effect in human HB in vitro is not much different from regular insulin. Therefore, these data suggest that insulin glargine is similar in mitogenic potency to AspB10 and IGF1, at least at low and presumably physiological concentrations.

Methylation of the cytosine C-5 position in the promoter region of tumor suppressor genes is an important mechanism of carcinogenesis in addition to gene mutation. However, the actual mechanisms of de novo methylation are not clear. We have reported the formation of 5-methylcytosine from cytosine in vitro, with methyl radicals generated from methionine sulfoxide (MetO). To confirm this reaction in vivo, MetO was added to the drinking water and administered to non-alcoholic steatohepatitis (NASH) mice, which develop hepatocarcinomas via endogenous oxidative stress. All of the animal experimental procedures were performed in accordance with the guidelines for the care and use of laboratory animals at Univ. Occup. Environ. Health. Histopathological examinations revealed incidences of hepatocellular carcinoma of 16.7% and 90% in the 0% and 3% MetO groups, respectively. Higher DNA methylation was detected in the promoter region of the p53 gene isolated from the livers of MetO-treated mice. The higher incidence of liver tumors may be due to the methyl radical-mediated formation of 5-methylcytosine in DNA, which triggers epigenetic changes. References: 1) Kawai K, et al. DNA methylation by dimethyl sulfoxide and methionine sulfoxide triggered by hydroxyl radical and implications for epigenetic modifications. Bioorg Med Chem Lett. 2010; 20: 260-5. 2) Kaiwa H, et al. DNA methylation at the C-5 position of cytosine by a methyl radical: a link between environmental agents and epigenetic change. Genes and Environment. 2011; 33: 61-5. 3) Kawai K, et al. Methionine sulfoxide stimulates hepatocarcinogenesis in non-alcoholic steatohepatitis (NASH) mouse: possible role of free radical-mediated DNA methylation. Genes and Environment. 2012; 34: 123-8.
1097 Deletion of Hepatocyte Nuclear Factor 4 Alpha Promotes Diethylamino-Lasoxinic-Induced Hepatocellular Carcinoma.

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HNF4α is known as the master regulator of hepatocyte differentiation. However, role of HNF4α in regulation of hepatocyte proliferation is not known. We investigated potential role of HNF4α in regulation of hepatocyte proliferation using a novel transmissible inducing hepatocyte specific HNF4α knockout mouse model. Hepatocyte specific deletion of HNF4α in adult mice resulted in increased hepatocyte proliferation with a significant increase in liver/body weight ratio. RNA sequencing-mediated global gene expression analysis revealed a significant number of the 500+ up-regulated genes are associated with cell proliferation and cancer. Further, a combined bioinformatic analysis of ChIP-sequencing and RNA-seq sequencing data indicated that a substantial number of up-regulated genes are putative HNF4α target genes. IPA-mediated functional analysis revealed the most significantly activated gene network after HNF4α deletion is regulated by c-Myc. To determine role of HNF4α in pathogenesis of HCC, we performed the classic initiation-promotion experiment using diethylamino-Lasoxinic (DEN). Deletion of HNF4α resulted in extensive promotion of DEN-induced hepatic tumors. HNF4α deletion resulted in 4-fold higher hepatic tumors, which were highly proliferative and less differentiated. Further, the HCC observed in HNF4α deleted mice exhibited significant up regulation of c-Myc and its target genes. These data indicate that HNF4α inhibits hepatocyte proliferation, is a potential tumor suppressor in the liver and plays a critical role in chemical carcinogenesis.

1098 EGCG Elicits Stage-Specific Sensitivity in a Novel Prostate Cancer Progression Model.

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Epigallocatechin gallate (EGCG), a major tea catechin, has been shown to have protective effects in a mouse model of PRCA, however, the relevance to human PRCA and the mechanism of action are less understood. We made the novel discovery that EGCG is a heat shock protein 90 (hsp90) inhibitor. Therefore, we utilized non-tumorigenic, tumorigenic, and metastatic PRCA cells from a human PRCA progression model to test the hypothesis that malignant cells are more sensitive to EGCG in a stage-specific manner and that sensitivity is related to the ability of EGCG to act as a hsp90 inhibitor. Treatment of cells with EGCG (25-50μM) in vitro decreased the viability and proliferation, and induced apoptosis selectively in PRCA cells compared to non-tumorigenic cells. EGCG also led to a decrease in the motility of tumorigenic PRCA cells, as analyzed by scratch assay, at the same concentrations tested. Moreover, when tumorigenic or metastatic cells were grown in vivo, mice supplemented with 0.06% EGCG in drinking water had significantly smaller tumors than those of the untreated group. To elucidate the mechanism of EGCG sensitivity, we performed affinity chromatography with EGCG-Sepharose. Binding assays revealed that hsp90 from metastatic cells had a higher affinity for EGCG than non-tumorigenic cells. Furthermore, EGCG disrupted hsp90 complex formation, as analyzed by Native PAGE, leading to an accumulation of dimeric hsp90, and promoted the degradation of hsp90-client proteins such as HER2, p-Akt and Raf1 in metastatic cells at concentrations that affected viability. These data suggest that EGCG may be an efficacious small molecule for the treatment of PRCA because it selectively targets PRCA cells and inhibits a molecular chaperone involved in many pro-cancer signaling cascades. Future studies will test if analogs of EGCG are more stable/bioavailable, and therefore more potent hsp90 inhibitors, for the treatment of PRCA. Funded by NIH Grant AT006366.

1099 Fluopyram: Mechanistic Investigations to Elucidate the Moa for Thyroid Tumor Formation in the Mouse.

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Fluopyram, a broad spectrum fungicide, caused an increased incidence of thyroid follicular cell (TFC) adenomas in males at the highest dose evaluated (750 ppm equating to 105 mg/kg/d) in the mouse oncogenicity study. Mechanistic studies were conducted on the male mouse to characterize the mode of action (MoA) for the thyroid tumor formation and to determine if thresholds exist for each key event. The proposed MoA consists of an initial effect on the liver by activating the constitutive androstane (CAR) and pregnane X (PXR) nuclear receptors causing increased elimination of thyroid hormones followed by an increased secretion of thyroid stimulating hormone (TSH). This change in TSH secretion results in an increase of TFC proliferation which leads to hyperplasia and eventually adenomas after chronic exposure. CAR/PXR nuclear receptors were shown to be activated from as early as 3 days of treatment as indicated by increased activity of specific Phase I enzymes (PROD and BROD respectively). Furthermore, evidence of increased T4 metabolism was provided by the induction of phase II enzymes known to preferentially use T4 as a substrate. Additional support for the proposed MoA was given by an increase of Tsh transcripts in the pituitary gland. Finally, increased TFC proliferation (Brdu incorporation) was observed after 28 days of treatment. In these dose response studies, clear thresholds were established for liver enzyme activities, T4 and TSH changes and TFC proliferation. Furthermore, each early change was shown to be reversible following fluopyram withdrawal. In conclusion, these studies indicate that fluopyram is a thyroid carcinogen and the resultant increased incidence of TFC adenoma in the male mouse is mediated by CAR/PXR activation, with subsequent TFC proliferation. Since liver mediated thyroid toxicity has been reported to be rodent specific, it is unlikely that fluopyram would induce thyroid changes and tumors in humans.

1100 Oncogene PKC Epsilon Inhibits Transcription Factor Nrf2 through Phosphorylation of Its Regulatory Molecule IκBα (Keap1).

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Nrf2/Keap1 (Keap1) serve as sensors of chemical and radiation stress and are critical in protection against oxidative stress and cellular transformation. IκBα functions as an adaptor protein for Cul3-Rbx1-mediated ubiquitination and degradation of Nrf2. Exposure to stressors leads to dissociation of Nrf2 from IκBα, stabilization and nuclear translocation of Nrf2 and activation of Nrf2 downstream cytoprotective proteins leading to cellular protection and survival. The molecular signal(s) that control Nrf2 control of Nrf2 remain elusive. In this report, we demonstrate that oncoprotein protein kinase C epsilon (PKCe) phosphorylates IκBα at amino acid positions Ser399 and Ser602 that is essential for IκBα/Nrf2 interaction and ubiquitination/degradation of Nrf2. Therefore, PKCe by regulating IκBα controls Nrf2 activation of downstream cytoprotective proteins and cell survival. PKCe phosphorylation of Nrf2 appears specific for recognizing Nrf2 since it does not affect Nrf2 interaction with other proteins including ubiquitin factors Cul3/Rbx1. Inhibition of PKCe or MEFs lacking PKCe or Nrf2S602A mutant all failed to phosphorylate IκBα and led to reduced degradation of Nrf2 and enhanced Nrf2 downstream signaling resulting in increased cell survival, proliferation and drug resistance. An analysis of protein microarray data from human lung and liver tumors showed lower PKCe and higher Nrf2. This inverse relationship between PKCe and Nrf2 in tumors presumably contributed to cancer cell survival and drug resistance. In conclusion, this is the first report that demonstrates a role of PKCe in phosphorylation of Nrf2 and control of Nrf2 with significant implications in survival of cancer cells that often express lower levels of PKCe.

1101 A Weight-of-Evidence Approach to Determine the Leydig Cell Tumor Mode-of-Action (MoA) for Pronamide.

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Pronamide, a herbicide, caused an increased incidence of Leydig cell tumors (LCT) in CD rats at 1000 ppm (27% vs. 9% in controls) and liver hypertrophy after two years of dietary exposure. There was no Leydig cell hyperplasia or tumors at the 12-month interim necropsy. Dose levels of 40 or 200 ppm had LCT incidences similar to controls, identifying a clear effect threshold. A series of studies was performed to elucidate the MoA for rat LCT and its relevance to humans. Pronamide was negative for androgen and estrogen receptor binding, aromatase inhibition, steroidogenesis alterations, and estrogen receptor transactivation. Despite no evidence of direct action upon the endocrine system, in vivo MoA studies revealed a slight, but consistent, increase in circulating levels of luteinizing hormone (LH), estradiol, and estrone in rats at 1000 ppm. It is proposed that the high-dose hormonal alterations are secondary to induction of liver-based testosterone metabolism/inactivation, coincident with liver hypertrophy and increased cytochrome P450. Secondary increases in LH are proposed to result in mitogenic stimulation of rat Leydig cells, which are quantitatively more sensitive than human Leydig cells to LH-driven proliferative response, based
primarily upon >10-fold more LH receptors per cell in rats vs. humans. This difference in sensitivity is supported by the >1000-fold difference between the incidence of human LCT vs. the rat. Overall, the weight-of-evidence for pronamide-induced rat LCT supports a threshold-response (i.e., nonlinear) approach for risk assessment due to the lack of genotoxicity, absence of direct endocrine effects, clear NOAEL for the LCT response (200 ppm), and an adequate margin of exposure for an effect in a uniquely sensitive animal model.

**1102** Haploregenotoxic Carcinogenic-Induced Temporal Changes in the Hepatic miRNAome.

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Non-genotoxic carcinogens are capable of inducing carcinogenesis via means other than direct modification of DNA structure. While there are many tests readily available for detecting genotoxic substances, methods for detecting and screening non-genotoxic chemical carcinogens are limited. Here we explore the potential of miRNA dysregulation as an indicator of non-genotoxic carcinogenesis by assessing the temporal-dependent changes in the miRNAome of Fisher rats. Ten chemical compounds, including: Phenobarbital (PB, 1000ppm), Chlordiazepoxide (ClAc, 1250ppm), Diethylyihexylphosphate (DEHP, 1200ppm), Monuron (Mon, 150ppm), Methapyrilene Hydrochlordide (MP HCI, 250ppm), 2-Aminoacetylfuorenone (genotoxic, 2-AAF, 40ppm), cinnamon antranilate (CA, 30,000ppm), Diethylyhxyadipate (DEHA, 25,000ppm), Benzo(b)fluorine (BbF, 1250ppm) and Diethylihthoureia (DETU, 250ppm), were fed to the diet in Fisher rats for a period of up to three months and liver samples were harvested at 7, 28 and 90 days. The total RNA was purified and analysed with Agilent Rat miRNA Microarrays Kit (Release 16.0). The expression of multiple hepatic miRNAs was found to be altered in response to chemical treatment in a temporally-dependent manner. Chemicals with a similar mode of action (e.g. DEHP and DEHA) induced similar responses on the hepatic miRNAome, confirmed by hierarchical clustering analysis ofqPCR. In the case of DEHA and DEHA, mir-101 family and mir-212 were transiently elevated (at least 3 fold) at 7 days of treatment. In contrast mir-200 family expression was unaffected at 7 days but was significantly induced at 90 days of treatment. Mir-101 family and mir-212 family are predicted to regulate MAPK pathway and target myc gene in the p53 feedback pathway, whereas the mir-200 family are involved in regulating the epithelial-mesenchymal transition and metastasis. These data suggest that miRNAs are potential biomarkers of non-genotoxic hepatocarcinogen and understanding the consequences of miRNA dysregulation aids the understanding of the mechanisms of carcinogenesis.

**1103** Dietary Carcinogenic PhIP induces Early DNA Damage and p53 Pathway Activation in CYP1A1-Humanized Transgenic Mice.

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2-Amino-1-methyl-6-phenylimidazo[4,5-b]pyridine (PhIP) is a dietary carcinogen and the most abundant heterocyclic amine generated from high temperature cooking of meat and fish. PhIP is activated via N2-hydroxylation to a proximate carcinogen primarily by cytochrome IIA2, CYP1A1-humanized (hCYP1A1) mice, which replaced the murine Cyp1a1 and Cyp1a2 with human CYP1A1 and CYP1A2, mimic human metabolism of PhIP. Previous study in our lab demonstrated that in hCYP1A1 mice, a single oral administration of PhIP induces high-grade prostatic intraepithelial neoplasia, mainly in the dorso-lateral glands (DLGs). Because murine DLGs correspond to the human prostate peripheral zone, the most common site of prostate cancer. PhIP-induced carcinogenesis in hCYP1A1 mice is a relevant mouse model for studying early stage prostate carcinogenesis. Herein, we investigated the early cellular and molecular events induced by PhIP in the dorso-lateral prostatic hCYP1A1 mice. Using immunohistochemistry, we detected strong positive staining of 8-Oxo-2′-deoxyguanosine as well as γ-H2AX in the DLGs on Days 1 and 3 after PhIP treatment, suggesting DNA damage and DNA strand breaks. Through gene expression profiling analysis and RT-qPCR, we found changes in expression of numerous p53-associated genes, including p21 and cyclin D1, indicating p53 activation and likely cell cycle arrest in PhIP-treated hCYP1A1 mice. Similar gene alterations were not observed in the PhIP-treated wild-type C57BL/6j mice. Altogether, these results suggest PhIP-induced DNA damage causing activation of the p53 pathways that may be involved in the early stage of carcinogenesis in the prostate. (Supported by NIEHS training grant ST32ES007148-25 and NIH grant RO1CA133021 as well as shared facilities funded by CA27720 and ES05022).

**1104** Reexamination of Initiator Dose in Ultra-Short-Term Carcinogenicity Study in RASH2 Mice.

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We have established an ultra-short-term carcinogenicity study in which the target organ was the skin, one of the common targets in rash2 mice. Although we previously reported that the suitable dose of 7,12-dimethylbenz[a]anthracene (DMBA) as an initiator was 50 mg at SOT 2010, several tumors were induced in DMBA treated groups without promotion, suggesting the dose of DMBA as an initiator was too high. In this study, we reexamined the suitable dose of DMBA as an initiator. Several compounds were applied to shaved dorsal skin of female rash2 mice one week after 12.5 or 25 μg DMBA application: oleic acid diethanolamine condensate (OADEC, 30 mg/kg, 7 times/week) or benzenethionium chloride (BC, 1.5 mg/kg, 5 times/week), known to have no promoting effect, 99.5% ethanol (7 times/week) or anhydroxy ethanol (5 times/week) as vehicles, or benzyl peroxide (BPO, 20 mg/head, 5 times/week) as a positive tumor promoter. Applied skin was studied histopathologically at eight weeks after DMBA application. Although no tumors were induced in OMDC, BC or vehicle treated groups, skin tumors were induced at five weeks after DMBA application in BPO treated groups, and the incidence of skin tumors reached 100% by seven weeks after DMBA application. The number of skin tumors, which were diagnosed as hyperplasia and/or papilloma, at necropsy were 29.2 and 35.6 in 12.5 μg and 25 μg DMBA treated groups, respectively. Therefore, we concluded that 12.5 μg DMBA was suitable to initiate the skin of rash2 mice in an ultra-short-term skin carcinogenicity study.

**1105** Modulation of Cocarcinogenic Effect of Aflatoxin B1 and Fumonisin B1 in a Short-Term Bioassay by Uniform Particle Size Novasil Clay.

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Co-contamination of AFB1 and F1 in corn-based food and feed is a health concern for their combinatorial toxic effects as well as potential co-carcinogenic effect. Uniform particle size Novasil (UPSN) clay has been used as an enterosorbent to reduce AFB1 and F1 exposure in animals and humans. In this study male F344 rats (150g) were randomly divided into five groups: negative control group, AFB1+F1 group, AFB1+low UPSN (0.25%) group, AFB1+F1+high UPSN (0.5%) group, and positive control group treated with diethylnitrosamine and 2-acetylaminoflourene. All treated animals were sacrificed after their last exposure. Liver was dissected and processed with regular H&E staining for examination of histological alterations. Liver tissue slides were also immunohistochemically stained for examination of formation of placent form glutathione S transferase positive (GST-P) foci. No detectable lesions were found in the negative control group, while the positive control group showed significant high rates of dysplasia (262 ± 29), prolific foci (53 ± 16), and nodules (15 ±10), as well as high rate of GST-P positive liver foci (58 ± 11), which demonstrates the success of the experiments. Co-exposure to AFB1 and F1 induced comparable rate of dysplasia (258 ± 40) to the positive control group and a higher rate of apoptosis (26 ± 8) than the positive control group (4 ± 4). UPSN in both low and high groups significantly inhibited formation of preneolesion in liver: For dysplasia, reduction of 46% (140 ± 29) and 56% (113 ± 20); for apoptosis, reduction of 42% (15 ± 4) and 57% (11 ± 4); for proliferative foci, reduction of 74% and 94%: for GST-P positive foci numbers, areas, and mean treated value of reduction was 93% and 94% and 31% and 53% were found, respectively. These results demonstrate significant modulation of UPSN on reducing potential co-carcinogenic effect in this short-term bioassay.

**1106** Validation of Cocarcinogenic Effects of Aflatoxin B1 and Fumonisin B1 in a Short-Term Bioassay.

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A short-term animal assay using Aflatoxin B1 (AFB1) as the initiator and fumonisin B1 (FB1) as the promoter demonstrated dramatic increase in formation of placental form glutathione S transferase positive (GST-P) foci in rat liver. In this
study, we validated this short-term bioassay and studied potential interactions be-
 tween AFB1 and BaP on the biochemical and histological changes and the forma-
tion of liver GST-P+ foci through a modified treatment protocol. AFB1 (150 μg) 
containing diet caused dramatic dysplastic hepatocytes, mild bile duct proliferation 
and necrosis, and proliferation foci formation. In contrast, BaP (250 ppm) con-
taining diet induced less dysplastic hepatocytes and more apoptotic bodies while no 
proliferation foci were found. Additive effects on increasing serum ALP, ALT, and 
AST activities were found in co-exposed animals as compared to single AFB1 or 
BaP treated. Sequential co-exposure to AFB1 and BaP led to significantly in-
creased numbers of dysplastic hepatocytes, apoptosis and proliferation foci. Co-ex-
posure of AFB1 and BaP also significantly increased the numbers of liver GTP-P+ 
foci by approximately 7.2 and 12.2 folds (p<0.01), and increased the mean sizes of 
GST-P+ foci by 6.0 and 7.5 folds (p<0.01), as compared to single AFB1 or BaP 
treated, suggesting strong synergistic effects in enhancing the preneoplastic lesion 
in liver. This study validates co-carcinogenic effects of AFB1 and BaP in the short-
term bioassay in rat liver and provides evidence for the use of this model in future 
intervention studies and mechanistic investigations.

1107 Tumor Promoter Action of Chromium-Containing Stainless 
Steel Welding Particulate Matter in the Lungs of A/J Mice. 
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Epidemiology studies show that occupational exposure to metal-rich welding par-
ticulate matter (PM) increases lung cancer risk. However, animal studies are lacking 
to conclusively link welding with an increased cancer risk. PM derived from stain-
less steel (SS) welding practices, in particular, contains carcinogenic metals such as 
hexavalent chromium and nickel. Previously, we found that PM derived from gas 
metal arc (GMA) welding of SS caused a borderline increase in lung tumor inci-
dence and a significant, chronic immune response in the lungs of mice. Thus, we 
assumed that PMd welding PM may act as a tumor promoter and increase lung 
tumor multiplicity. In this study, the capacity of GMA-SS welding PM to promote 
lung tumors was evaluated using a 2-stage (initiation-promotion) model in lung 
tumor susceptible A/J mice. Age and weight-matched male mice (n=26-29/group) 
were treated either with the initiator 3-methylcholanthrene (MCA;10 μg/gi.p.) or 
vehicle (corn oil; CO) followed by 5 weekly pharyngeal aspirations of GMA-SS (340 
or 680 μg) or PBS. Lung tumors were enumerated at 30 weeks following ini-
tiation with MCA. Body weights were recorded at 2 week intervals and no effect 
of treatment was found. MCA initiation followed by GMA-SS exposure promoted 
lung multiplicity in both the lower dose (12.0 ± 1.5 tumors/mouse; 
p=0.0001) and high dose (14.0 ± 1.8 tumors/mouse; 
p=0.0001) groups signifi-
cantly above that of MCA/PBS (4.77 ± 0.7 tumors/mouse). Not only was this 
highly significant for the average total number per mouse, multiplicity was also 
increased (p<0.004) across all live individual lung regions of GMA-SS-exposed mice. 
No treatment effects were found in the corn oil groups at 30 weeks and also, as 
expected, tumor incidence was greater than 93% in the MCA treated groups which 
confirmed our earlier results. In conclusion, GMA-SS welding PM is a lung 
tumor promoter in vivo. These novel findings implicate that susceptible indi-
viduals may be at greater risk for lung cancer development after exposure to weld-
ning PM.

1108 Induction of 1, 3-Dichloropropene Rat Liver Tumors 
through a Nongenotoxic Mode of Action. 
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1,3-Dichloropropene (1,3-D) is a soil fumigant used for control of parasitic nema-
todes. It was previously shown that dietary exposure to 1,3-D was associated with 
an increased incidence of benign hepatocellular adenomas in male rats in one of 
two rat cancer studies. In in vivo genotoxicity studies, 1,3-D was shown consis-
tently to be non-mutagenic in studies where physiologic levels of glutathione and 
glutathione-S-transferases were present and genotoxic stabilizers were not present, 
thus supporting liver tumorigenesis through a non-genotoxic mode-of-action 
(MoA). The present study was undertaken to investigate the hypothesis that 1,3-D 
induces rat liver tumors by acting as a tumor promoter in the liver carcinogenesis 
process. For assay and potential 1,3-D tumor promotion, dityramine-induced 
fischer 344 rats were treated by gavage with 25 mg/kg/day 1,3-D or 80 mg/kg/day 
phenobarbital (PB) for either 30 or 60 days, or for 30 days followed by 30 days recov-
ery. Staining for the placental form of glutathione S-transferase (GSTP) was 
documented post-exposure as a phenotypic marker of preneoplastic foci. While hav-
ing no effect on GSTP positive foci, 1,3-D significantly enhanced the number and 
size of GSTP negative lesions at 30 and 60 days. Importantly, after 30 days recov-
ery, the number and size of GSTP negative lesions in 1,3-D treated animals re-
turned to control levels thus demonstrating 1,3-D promotion of foci and not 
tumor initiation. Also supportive of tumor promotion, the hepatocyte labeling 
index (LI) in GST-P negative foci was increased in 1,3-D treated rats at 30 and 60 days and returned to control levels after 30 days recovery. PB, a well recognized non-genotoxic rodent liver carcinogen, 
produced an expected reversible increase in number and size of the GSTP positive lesions. The results pre-
sent are consistent with 1,3-D inducing liver carcinogenesis through a non-geno-
toxic MoA by induction of reversible proliferation of non-GSTP staining focal 
populations of hepatocytes.

1109 Effects of Long-Term Administration of the Tissue-Selective 
Estrogen Receptor Modulator Bazedoxifene on Survival and 
Tumor Formation in Rats. 
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2Abbott Laboratories, Abbott Park, IL; 3Pharmacokinetics, Dynamics, and 
Metabolism, Pfizer, La Jolla, CA; 4Medical Affairs, Pfizer, Collegeville, PA; 5Division 
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Bazedoxifene acetate (BZA) is a selective estrogen receptor modulator that is 
approved in a number of countries for the prevention and/or treatment of osteopor-
osis in postmenopausal women. To assess for carcinogenic potential, BZA was ad-
ministered ad libitum in the diet to male and female rats for 2 years. BZA resulted 
in a reduction and a delayed onset in total tumor burden in both male and female 
rats. Survival rates were enhanced due to decreased pituitary (males and females) 
and mammary tumors (females) and decreased body weight gain in BZA-treated 
animals compared to controls. BZA caused an increased incidence of benign ovar-
ian tumors and renal tubular tumors (males). Results from separate studies sug-
gested that BZA caused an increase in LH, which stimulated formation and persist-
ence of ovarian cysts, which eventually progressed into benign ovarian granulosa 
cell tumors. The reduction in pituitary and mammary gland tumors were attrib-
uted to BZA-related antagonism of endogenous estrogens at the estrogen receptors. 
The greater increase in renal tumor incidence in male rats given BZA was associated 
with the increased survival and increased time for development of late onset tu-
mors. These findings are consistent with a non-genotoxic mechanism, unique to 
male rats, that involves test article-induced corticomedullary mineralization, renal 
tubular and stromal proliferation, and exacerbation of naturally occurring chroni-
CIC nephropathy in aged male rats that leads to proliferative changes and tumor forma-
tion. In conclusion, BZA elicited agonistic or antagonistic effects in a tissue-selec-
tive manner which was generally consistent with expression and/or activity of estro-
gen receptors in those tissues.

1110 Resveratrol Exposure Reduces Benzo(a)pyrene-DNA Adduct 
Concentrations in Apcmin Mouse Model of Colon Cancer. 
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Colon cancer is the third leading cause of cancer cases and related deaths as per the 
statistics provided by the American Cancer Society. Exposure to toxins such as 
benzo(a)pyrene (BaP) is one of the contributing factors to the development of spor-
adic colon cancer. Our studies have shown a decrease in the size, and number of 
adenomas in the colon of mice exposed to BaP and resveratrol (RVT), compared to 
BaP exposure alone. We have also shown that RVT exposure caused a decrease in 
the expression and activity of CYP1A1/B1 enzymes and BaP metabolite concen-
trations both in liver and colon. Since DNA damage is one of the key events in car-
inogenesis, the objective of this study was to investigate whether RVT exposure si-
multaneously or prior to BaP treatment alters BaP-DNA adduct concentrations in 
Apcmin mouse. The treatment consisted of BaP only administration (in peanut oil) 
at a dose of 0.1 μg/kg bw via gavage over a 4-day period (group I); BaP (100 
μg/kg bw) co-administered with RVT (in 10% ethanol + 90% deionized water) at 
a dose of 45 μg/kg bw (group II); RVT administered for 1 week prior to BaP dos-
ing (group III). Post exposure, DNA was isolated from colon and liver samples, and 
BaP-DNA adducts were determined using a radioactively labeled adduct detection 
method. The adduct concentrations showed a trend that mirrored the concentrations of BaP metabolites in the organs studied with low concentrations in 
RVT-treated mice compared to BaP-treated mice. Between the two RVT-treatment 
strategies, concurrent administration of RVT appeared to affect the BaP bioactiva-
tion compared to RVT treatment prior to BaP exposure as reflected by the low 
adduct concentrations in the former group III compared to groups I & II. Overall, 
our results suggest that RVT provides a preventive effect against BaP-induced colon 
cancer progression in Apcmin mouse (funded by NIH grants 5F31ES019432-01A1, 
5R01CA142845-02, 5T32HL007735-12, and 5R25GM059994-11).

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1111 Triclosan Promotes the Development of Hepatocellular Carcinoma in Mice.

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Triclosan (TCS), a chlorophenol, is used in a large number of personal care products as an antibacterial agent. TCS has endocrine disruption properties, has been detected in microgram per liter levels in various water ways in the US, and is considered to be a major environmental contaminant in aquatic ecosystems. Daily exposure to TCS has resulted in its detection in humans with studies showing its presence in plasma, breast milk, and urine. Studies have increasingly linked TCS to a range of health and environmental effects; however, there are no mechanism-based studies that directly address TCS exposure to negative health effects in humans. Using wild type and Car/-/- mice, TCS has been shown to effectively induce hepatic Cyp2b10 in a Car-dependent manner. However, transient transfection experiments revealed that TCS is not a direct CAR agonist but behaves as a phenobarbital-like inducer that indirectly activates CAR and facilitates its translocation to the nucleus. Using the chemical procarcinogen diethylnitrosamine (DEN) to initiate tumorigenic episodes in mice, TCS exposure through drinking water promotes the development of hepatocellular carcinoma (HCC). The development and promotion of HCC by TCS, compared to DEN-only treated mice proceeds in a CAR-dependent fashion. This finding stands in contrast to CAR-dependent tumor induction with the liver tumor promoter phenobarbital. HCC tumors produced by DEN-TCS exposure lead to a number of changes in the liver including histological alteration in hepatocytes, increases in liver-to-body weights, elevated levels of α-fetoprotein, and enhanced levels of tumor-promoting cytokines IL-6 and TNFα. In addition, tumor-bearing livers significantly alter the expression profile for genes associated with drug metabolism. The identification of TCS as a liver tumor promoter may be significant because of its abundance and long term use in consumer products and its propensity for rapid absorption into the systemic circulation. (Supported by USPHS grant ES010357)

1112 Kupffer Cells Modulate Wyeth-14, 643 and Phenobarbital-Induced Hepatocyte Proliferation in Naïve and Diethylnitrosamine (DEN)-Initiated Mice.

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Kupffer cells (KC) play an important role in liver homeostasis. Upon activation, KC release inflammatory and growth regulatory mediators that have been linked to acute and chronic liver responses including hepatic cancer. Wyeth 14,643 (WY) and Phenobarbital (PB) are non DNA reactive hepatic carcinogens in rodents. The present study was conducted to investigate the role of KC in liver cell proliferation in naïve and focal lesion containing mice. In our initial study, KCs in male C3H mice were depleted using liposome encapsulated clodronate (Lip-cld, 0.1 ml/10 g, iv). LPS, a known KC activator, was used as a positive control (1 mg/kg, ip, 2X/wk). Lip-cld treatment resulted in depletion of 99% of KC as confirmed by F4/80 positive staining cells. WY treatment increased the relative liver weight and DNA synthesis in naïve mice after 7 and 21 d treatment. LPS and PB increased hepatic DNA synthesis (7 and 21 d). Following KC depletion, the induction of DNA synthesis by LPS and PB were significantly decreased, while WY-induced DNA synthesis was not altered. In the second study, mice were injected with DEN (50 mg/kg, ip, 2X/wk) (4 wk). Following development of focal lesions, mice were treated with WY or PB for 28 d. Treatment with PB or WY increased the relative focal volume and depletion of the KC decreased the relative focal volume. WY and PB increased DNA synthesis in both basophilic and eosinophilic lesions, similar to LPS. Upon KC depletion, a decrease in DNA synthesis was seen in PB treated mice. In the surrounding non-lesion tissue, PB produced a marginal increase in focal volume and depletion of the KC decreased the relative focal volume. WY and PB treated with WY or PB for 28 d. Treatment with PB or WY increased the relative focal volume and depletion of the KC decreased the relative focal volume. WY and PB were significantly decreased, while WY-induced DNA synthesis was not altered. In the second study, mice were injected with DEN (50 mg/kg, ip, 2X/wk) (4 wk). Following development of focal lesions, mice were treated with WY or PB for 28 d. Treatment with WY or PB increased the relative focal volume and depletion of the KC decreased the relative focal volume. WY and PB were significantly decreased, while WY-induced DNA synthesis was not altered. In the second study, mice were injected with DEN (50 mg/kg, ip, 2X/wk) (4 wk). Following development of focal lesions, mice were treated with WY or PB for 28 d. Treatment with WY or PB increased the relative focal volume and depletion of the KC decreased the relative focal volume. WY and PB were significantly decreased, while WY-induced DNA synthesis was not altered. In the second study, mice were injected with DEN (50 mg/kg, ip, 2X/wk) (4 wk). Following development of focal lesions, mice were treated with WY or PB for 28 d. Treatment with WY or PB increased the relative focal volume and depletion of the KC decreased the relative focal volume. WY and PB were significantly decreased, while WY-induced DNA synthesis was not altered. In the second study, mice were injected with DEN (50 mg/kg, ip, 2X/wk) (4 wk). Following development of focal lesions, mice were treated with WY or PB for 28 d. Treatment with WY or PB increased the relative focal volume and depletion of the KC decreased the relative focal volume. WY and PB were significantly decreased, while WY-induced DNA synthesis was not altered. In the second study, mice were injected with DEN (50 mg/kg, ip, 2X/wk) (4 wk). Following development of focal lesions, mice were treated with WY or PB for 28 d. Treatment with WY or PB increased the relative focal volume and depletion of the KC decreased the relative focal volume. WY and PB were significantly decreased, while WY-induced DNA synthesis was not altered.

1113 Bioassay-Directed Fractionation of Diesel and Biodiesel Emissions.

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Biofuels are being developed as alternatives to petroleum-derived products, but published research is contradictory regarding the mutagenic activity of such emissions relative to those from petroleum diesel. We performed bioassay-directed fractionation and analyzed the polycyclic aromatic hydrocarbon (PAH) levels of particles generated using petroleum diesel (B0) and those from soy-based biodiesel where the soy accounted for 20, 50 or 100% of the fuel (B20, B50, B100) from a Yanmar L70 diesel engine. We also evaluated a composite sample of diesel-exhaust particles (C-DEP) generated from petroleum diesel with a 30-kW 4-cylinder Deutz BF4M1008 diesel engine connected to an air compressor. The biodiesel and C-DEP particles were extracted with dichloromethane (DCM), and the percentage of extractable organic material of these complex environmental mixtures were determined. The extracts were then solvent exchanged into dimethyl sulfoxide and evaluated for mutagenicity in various Salmonella strains +/-S9. We calculated the mutagenic emission factors (revertants/megajoule, rev/MJ) from data from strain TA100 +/-S9, which responds to PAH-type mutagenicity. These mutagenic emission factors for the biodiesels were 3X less (B20), 6X less (B50), and 8X less (B100) than that of petroleum diesel (B0). The whole extract of C-DEP (petroleum diesel) was sequentially fractionated with solvents of increasing polarity, and >50% of the mass eluted in fraction 1; this fraction was not mutagenic. The 2nd fraction had 60% of the TA98-S9 activity, indicative of nitrosamines. We conclude that under these experimental conditions, emissions from biodiesel were less mutagenic than those from petroleum diesel, and based on the fractionation of the C-DEP, most of the activity is associated with PAHs. [Abstract does not necessarily reflect the views or policies of the U.S. EPA.]


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CBF-g-box binding factor-A is a member of the heterogeneous nuclear ribonucleoprotein (hnRNPs) family of RNA-binding proteins involved in a variety of cellular functions, including transcriptional regulation. We showed that CBF-A (hnRNPN A/B) regulates the expression of the Ha-ras oncogene by binding to enhancer elements (ets-like) in the promoter region. Deregulated expression of CBF-A isoforms is associated with mammary carcinogenesis in NMu-tREATED rats. CBF-A is a regulator of epithelial-mesenchymal transition (EMT), an important process in the cancer progression. To evaluate the role of CBF-A, we generated knockout mice. Behavioral evaluation in the elevated plus maze showed that CBF-A null mice spent ~300% more time in the open arms of the maze compared to the wild-type, suggesting that loss of CBF-A results in a low-anxiety behavioral phenotype. This behavioral abnormality may be related to the role of CBF-A in regulating the expression of vasopressin in the brain. To further study the role of CBF-A in tumorigenesis, we exposed wild-type mice, as well as CBF-A null and heterozygous mice, to NMU to initiate the development of the mammary gland. DNA-methylating agent ENU (100 μg/mL at 9-10 weeks of age). While administration of ENU decreased the survival rate in all genotypes compared to vehicle control mice, CBF-A null mice had a decreased survival rate and tumor-free survival rate compared to the wild-type. Taken in concert, these data suggest that CBF-A plays an important role in maintaining both normal growth control and behavior. (Supported by Grants from the DOD (BC971045), the NIEHS funded Toxicogenomics Research Consortium (EO111387), NIEHS Center Grants (ES050522 and ES007033) and CNIN (CA072720).

1115 Inhibition of Hepatocyte Nuclear Factor 1 and 4 Alpha (HNF1α and HNF4α) as a Mechanism of Arsenic Carcinogenesis.

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Inorganic arsenic (i-As) is a naturally occurring toxic metalloid affecting millions of people worldwide. It is known to be carcinogenic, liver being a potential target, and related to the prevalence of diabetes in arseniasis-endemic areas. Hepatocyte nuclear factor 1 and 4 alpha (HNF1α and HNF4α) are key members of a transcriptional network essential for normal liver architecture. Changes in HNF1α and HNF4α expression are clearly associated with the development of liver malignancies and diabetes in humans. In this work, hepatic HepG2 cells and Golden Syrian hamsters were exposed to sub-toxic, environmentally relevant doses of sodium arsenite (5A; up to 10 μM in vitro, 15 mg/L in vivo) in order to evaluate whether arsenic is able to compromise the expression of hepatocyte nuclear factors (HNFs). Also, liver histopathological examination was carried out and several markers of hepatocyte dif-
ferentiation and glucose metabolism status were determined as a measure of i-As-induced toxicity/apoptosis, (ii) attained loss of tissue specific features (as shown by the observed downregulation of ALDOB, PEPCK and CYP1A2, the triggering of the epithelial-to-mesenchymal transition program (EMT) and the hyper-secretion of matrix metalloproteinase-2 and 9), (iii) failed to maintain balanced the expression of the "stemness" genes C-MYC, OCT3/4, LIN28 and NOTCH2 and (iv) showed glucose metabolism impairment. We conclude that the i-As-induced down-regulation of HNF1α and HNF4α under chronic settings may play a central role in the features of disease and cancer observed both in vivo and in vitro.

**1116 Carcinogenicity of Diesel (DEE) and Gasoline Engine Exhaus (GEE).**


Recently, an IARC Monographs Working Group (WG) reevaluated the carcinogenic hazards to humans of DEE and GEE. DEE was classified as "carcinogenic to humans" (Group 1) and GEE as "possibly carcinogenic to humans" (Group 2B). A US study in non-metal miners included a cohort analysis and a nested case-control analysis adjusted for tobacco smoking. Both showed positive trends in lung cancer risk with increasing exposure, with a 2.3-fold increased risk in the highest category of cumulative or average exposure. A 40% increased risk for lung cancer was observed in U.S. railroad workers exposed to DEE. Indirect adjustment for smoking and a more accurate exposure assessment strengthened the validity of the results. A large cohort study reported a 15–40% increased lung cancer risk in US truck drivers and dockworkers with exposure to DEE, with a significant trend of increasing risks with longer duration of employment, adjusted for tobacco smoking. Findings in other occupational groups and case-control studies including various occupations with similar exposures supported the WG's conclusion of "sufficient evidence" in humans for the carcinogenicity of DEE. The WG concluded that there was "sufficient evidence" in experimental animals for the carcinogenicity of whole DEE, DEE particles, and extracts of the particles, which also induced, in vitro and in vivo, various forms of DNA damage. Positive genotoxicity biomarkers of exposure and effect were also observed in humans exposed to DEE. The WG concluded that there is "strong evidence" for the ability of whole DEE to induce cancer in humans through genotoxicity. GEE and cancer risk was investigated in only a few epidemiological studies and, because of the difficulty to separate the effect of DEE and GEE, evidence for carcinogenicity of GEE was evaluated as "inadequate". Organic extracts of GEE condensate induced a cancer in rats. The WG concluded that there was "sufficient evidence" in experimental animals for the carcinogenicity of whole DEE, DEE particles, and extracts of the particles, which also induced, in vitro and in vivo, various forms of DNA damage. Positive genotoxicity biomarkers of exposure and effect were also observed in humans exposed to DEE. The WG concluded that there is "strong evidence" for the ability of whole DEE to induce cancer in humans through genotoxicity. GEE and cancer risk was investigated in only a few epidemiological studies and, because of the difficulty to separate the effect of DEE and GEE, evidence for carcinogenicity of GEE was evaluated as "inadequate". Organic extracts of GEE condensate induced a cancer in rats. The WG concluded that there was "sufficient evidence" in experimental animals for the carcinogenicity of condensates of GEE.

**1117 Suicide Inhibition Is Responsible for the Paradoxical Absence of CYP2B10 Enzyme Activity in Mouse Liver following Nitropyrrin- or Pronamide-Induced CAR Activation.**

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Establishing a model of action (MoA) for a toxicological finding requires the identification of key events (both causal and associative) that progress to the apical point of interest. In the case of constitutive androstane receptor (CAR)-mediated rodent hepatocarcinogenesis, the first key event is activation of the CAR signaling pathway, which is often measured via the empirically observable biomarker of Cyp2b10 induction (transcript, protein, and/or enzyme activity). The agonistic promavide (herbicide) and nitrapyrin (nitrification inhibitor) induced hepato-cellular tumors in mice as a result of high-dose, long-term dietary administration. Subsequent MoA studies for each chemical were consistent with a role for CAR activation in the mouse liver tumorigenesis. With both chemicals significant increases in Cyp2b10 transcript as well as protein were observed; however, there was no associated change in the activity of the enzyme as measured by PROD. To investigate the role for suicide inhibition in this paradoxical finding, in vitro experiments were conducted using phenobarbital (PB)-induced microsomes. These microsomes were treated with PB (negative control), curcumin (positive control), nitrapyrin, or pronamide. In this system, nitrapyrin and pronamide inhibited PROD activity in a dose-related manner and up to 97% and 56% at 500 μM, respectively. While PB had no effect on PROD activity, curcumin had a dose-related inhibition of Cyp2b-mediated PROD activity of up to 63% at 40 μM. These results indicate that the two agrochemicals and their nitro derivatives irreversibly inhibited Cyp2b10-mediated PROD activity and elucidate the apparent inconsistency between protein levels and enzyme activity of treated livers while supporting the CAR-mediated MoA for liver tumorogenesis. Further, these data indicate that measurement of transcript levels as biomarkers for CAR activation is more appropriate than traditional enzyme activity (e.g., PROD) measurements.

**1118 Structure-Dependent Effect of Diindolylmethane Derivatives on Inactivation of the Oncogenic NRA1/NRA2 Receptor in Colon Cancer Cells.**

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The nuclear receptor 4A (NRA4A) subfamily members TR3 (NRA4A1) and Nurrl (NRA2) have been investigated in this laboratory as novel drug targets for clinical treatment of pancreatic cancer. Inactivating of these oncogenic transcription factors by analogs derived from the chemopreventive phytochemical diindolylmethane (DIM) exhibit promising in vitro and in vivo chemotherapeutic effects. Knockdown of NRA4A1 and NRA2 by RNA interference in colon cancer cells decreased cell proliferation, induced apoptosis and inhibited colon cancer cell invasion. Using NR4A4-responsive promoter elements (monomer-binding NBRE and dimer-binding NurRE) linked to a luciferase reporter gene, we investigated the structure-dependent activity of over forty 1,1-bis(3-indolyl)-1-(substituted)phenyl)methane (DIM-C-Phe) and 1,1-bis(3-indolyl)-1-(heteroaromatic)carbon (DIM-C-heteroaromatic) analogs as inactivators of NRA4A1, NRA2 or both receptors. In colon cancer cells we identified DIM-C-Phe analogs containing para-hydroxy, chloro, methyl, or trifluoroethoxy groups that inactivated NRA4A1 and the 3-pyridine analog (DIM-C-Pyr-3) inactivated NRA2-mediated transactivation. In contrast, several other p-substituted phenyl analogs containing fluoro, iodo or cyano groups were potent inactivators of both receptors. The mechanisms of NRA4A inactivation by C-DIMs were investigated and the possible role of site-specific phosphorylation was determined using several protein kinase inhibitors including LY294002, PD98059 and SP600125. The results show that in RKO colon cancer cells transfected with NBRE-luciferase construct and treated with DIM-C-Phe, NRA4A inactivation was inhibited by PD98059. These results indicate that the chemotherapeutic properties of C-DIMs are outcomes of NRA4A inactivation and the NRA1/NRA2-dependent pathways and role of kinases are currently being investigated.

**1119 Evaluation of microRNAs in Blood for Detection of Chemical-Induced Carcinogenesis.**

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MicroRNAs (miRNAs) are emerging as a valuable tool in toxicological applications due to their role in regulation of gene expression in different biological pathways. Multiple studies have revealed that miRNAs are present and relatively stable in clinically accessible biofluids such as blood. miRNAs in biofluids may provide a non-invasive way of detecting chemical carcinogenicity. In this study, we evaluated whether microRNA profiling and individual miRNAs in mouse blood can predict carcinogenesis induced by N-ethyl-N-nitrosourea (ENU). Male B6C3F1 mice were treated with either a single dose of 140 mg/kg ENU or vehicle alone. Blood was collected on post-treatment days (PTDs) 1, 2, 3, 7, 14, 28, and 42. Profiling of total miRNAs in the blood showed that a number of miRNAs associated with carcinogenesis were altered by the ENU treatment. The temporal alteration of one of these miRNAs, miR-34a, a tumor suppressor, by ENU was characterized further. The level of miR-34a expression was significantly increased by ENU as early as PTD 1, reached a maximum level of about 10-fold over controls at PTD 7, and then decreased to control levels at PTD 42. Serum levels of miR-34a were also examined to assess the contribution of blood cells to miR-34a levels in whole blood. The results suggest that miRNAs have potential use as early sensitive circulating biomarkers of carcinogenesis exposure.

**1120 Hemodynamic Changes, Arrhythmias, and Sudden Death in Dogs following Repeated Dosing of AKT/pp70S6K Inhibitor.**

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PF-0476340 (PF) is an inhibitor of AKT & pp70S6K. Target organs in GRP repeat dose studies in dogs included eye, GI, liver, testes. In addition, unexplained deaths were observed in a 10d dose range-finding study (3, 10, 30 mpk/d), mor-
Mechanisms that underlie the strong association between air pollution exposure and increased cardiovascular disease remain unknown. We hypothesize that soluble components of ultrafine particles (soluble UF) initiate procoagulant activities in endothelial cells following exposure to soluble UF. There was an immediate induction of TF expression in HCAEC following exposure to soluble UF. There was an immediate induction of extracellular and intracellular H2O2 production following soluble UF exposure, and treatment with antioxidants attenuated the UF-induced upregulation of TF, linking the procoagulant response to ROS formation. Multiple chemical inhibitors indicated NOX as the source of increased ROS, and by the finding that NOX-4 siRNA prevented UF-induced upregulation of TF mRNA. These data show that exposure to soluble UF induces endothelial procoagulant activity, which requires new TF synthesis through ROS production and by NOX-4. These novel findings provide mechanistic insights into the enhanced thrombosis and endothelial dysfunction associated with air pollution exposure and increased risk for CV morbidity and mortality.

1123 Evaluation of Age and Sex Effects on Mitochondria-Related Gene Expression in Rat Heart.

Age-related susceptibilities and sex-based differences in cardiovascular diseases are known and the underlying mechanisms need further investigation. Mitochondria are critical for normal cardiac function so alterations in such at different ages and sexes may affect the way the heart responds to drugs. To understand the role of mitochondria in age- and sex-related differences in the heart, we investigated the expression levels of mitochondria-related genes. Gene expression in the heart of male and female Fischer 344 rats at 8 (young), 21 (adult) and 78 (old) weeks of age was measured using Agilent whole genome rat arrays. Out of 18,435 unique genes on the Agilent rat microarray, 869 unique genes were determined to be mitochondria-related. Age effect was evaluated by using ANOVA coupled with pair-wise t-test (p<0.05) between each age group, and sex difference was evaluated at each age using t-test (p<0.05). The expression levels of genes involved in oxidative phosphorylation (Ox Phos), membrane transporters, and fatty acid (FA) metabolism were the highest at 21 wks compared to 8 or 78 wks in both males and females. A significant age-related effect was observed in 21% and 7% of Ox Phos genes and 15% and 7% of membrane transporters in males and females, respectively, at all age groups. Only male rats showed significant age-related effects on the expression levels of genes (21%) involved in FA metabolism. Significant sex-based differences were observed in the expression levels of genes involved in Ox Phos at 78 wks (19%); higher expression levels were noted in female rats compared to males. Membrane transporters showed significant sex differences at 8 wks (21%), 21 wks (40%) and 78 wks (13%). Genes involved in FA metabolism also showed sex-based differences at 8 (34%), 21 wks (29%) and 78wks (10%); most of the genes had higher expression in females than males. These findings may provide important insights into understanding the role of mitochondria in cardiovascular diseases in terms of sex based differences or altered susceptibility with age.

1124 Assessment of Postmarket Cardiac Adverse Events in Patients Treated with Antipsychotic Drugs.

This study assessed the cardiac adverse events (cAEs) induced by antipsychotic drugs, and evaluated their relationship to hERG affinities, clinical drug exposure and the magnitude of clinical QTc prolongation. Data were collected from public regulatory documents (e.g. Summary Basis for Approvals) and supplemented with published literature data. The incidence of each cardiac toxicity endpoint was calculated using the total number of all adverse events reported for each drug as a denominator. The incidence of QT prolongation was 5.1, 3.4, 0.5, 1.0, 0.9 and 1.9% for thioridazine, ziprasidone, quetiapine, risperidone, olanzapine, and haloperidol, respectively, which were correlated to the magnitude of QTc change from baseline (dQTc) of 40.7, 26.4, 19.5, 15.8, 12.7 and 11.3 msec, respectively. The incidence of Torsade de pointes (TdP) were 0.9, 0.7, 0.1, 0.2 and 0.1% for thioridazine, ziprasidone, quetiapine, risperidone, olanzapine, and haloperidol, respectively, which were correlated to the magnitude of QTc change from baseline (dQTc). Furthermore, the incidence of QT prolongation was correlated to the ratio of total plasma Cmax/hERG IC50 for the same group of drugs. Pimozide, one of the most potent hERG blockers (hERG IC50 > 18 nM), had the highest incidences in causing QT prolongation (7.2%), TdP (3.4%), long QT syndrome (0.7%), syncope (3.4%) and sudden cardiac death (0.3%). The incidences of ventricular arrhythmias and cardiac arrest were greater than equal to 5% in patients receiving dizapam, imipramine, thioridazine, haloperidol, amitriptyline, pimozide, or trazodone. The incidences of sudden death were more than 1% in pa-
patients treated with desipramine, thioridazine, ziprasidone or pimozide. In conclusion, relatively high incidences of potentially serious cardiac side effects, including QT prolongation, TdP, and lethal outcomes occurred in patients treated with the antipsychotics. The hERG affinities combined with the levels of clinical drug exposure may serve as one of the predictors for drug-induced cardiac toxicity by antipsychotics.

**1125 Ozone Coexposure Modifies Cardiac Function Responses to Fine and Ultrafine Particulate Matter in Mice.**

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There is growing evidence from epidemiological studies that show acute exposure to particulate matter (PM) increases the risk of cardiovascular morbidity and mortality. Although the data supporting these findings are increasingly more convincing, the immediate impact of PM inhalation on cardiac function needs to be further clarified; this is particularly true of multipollutant exposures. Thus, this study was designed to evaluate the cardiac effects of concentrated ambient fine (PM2.5) and ultrafine (UFP) particles with and without ozone (O3) co-exposure. Based on previous findings, we hypothesized that UFP would cause the greatest decrement in cardiac function and that O3 co-exposure would worsen the response. Mice were exposed by whole-body inhalation to either 200 µg/m3 PM2.5 or 100 µg/m3 UFP with or without 0.3 ppm O3; separate groups were exposed to either filtered air or O3 only. Twenty-four hours after exposure, cardiac function was assessed using a Langendorff cardiac perfusion preparation. Coronary flow, left ventricular developed pressure (LVDP) and contractility were measured before and after cardiac ischemia/reperfusion (I/R) injury. PM2.5 or O3 alone, or co-exposure to UFP+O3 caused a significant decrease in baseline LVDPD and contractility. Interestingly, UFP alone or PM2.5+O3 did not cause significant decrements in cardiac function when compared to controls, whereas there were significant differences in recovery LVDPD or contractility between any group after I/R injury. These data suggest that the cardiac effects of PM inhalation are dependent on particle size and that O3 interacts with PM2.5 and UFP differently, resulting in varied cardiac impacts. Thus, these findings indicate that the microvascular effects of particles and gas co-exposures are not simply additive or generalizable, which increases the complexity of risk assessment. (This abstract does not reflect EPA policy)

**1126 An Integrative Approach to Identify Functional, Structural and Pathological Biomarkers of Doxorubicin-Induced Cardiotoxicity.**

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Doxorubicin is an anthracycline antibiotic commonly used in treatment regimens for a wide range of malignancies, including hematological cancers, many types of carcinoma and soft tissue sarcomas. Cardiotoxicity is a major safety complication associated with the clinical use of doxorubicin, several other marketed anti-cancer drugs and with many pre-clinical drug candidates. There is a need for novel, translational biomarkers to aid cardiac structural and functional monitoring in both patients at risk of cardiotoxicity and in pre-clinical species administered early drug candidates. In the present study, a detailed characterization of the effects of chronic doxorubicin treatment on rat heart structure and function has been performed using a range of approaches. A rat model of doxorubicin-induced cardiomyopathy was developed, involving weekly intravenous boluses of doxorubicin hydrochloride followed by a 4 week ‘washout’ period. A time-course assessment of cardiac function using multiple MRI biomarkers was performed under recovery anaesthesia, prior to dosing, on days 1 (baseline), 15, 29, 43, 57 and under terminal anaesthesia prior to necropsy on day 78. Reductions in cardiac output and ejection fraction were observed in treated animals from day 15. Peak myocardial enhancement was observed at day 57 onwards and correlated with serum cardiac troponin I elevations. A detailed immunohistochemical cardiac assessment is currently being performed to provide an understanding of the relationship between the development of pathological and functional MRI changes, serological biomarker elevations and cardiac pathological lesions. The ultimate aim is to apply the learning to pre-clinical drug discovery in order to support safety validation of drug targets, to optimize chemical design and to ultimately progress the safest molecules into later pre-clinical and clinical testing.

**1127 Both Vitamin D Excess and Deficiency Increase Blood Pressure, but Only Excess Increases Arterial Stiffness.**

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Vitamin D deficiency has been linked to an increased risk of hypertension, myocardial infarction and stroke. Recently, the Tolerable Upper Intake Level of 1, 25-dihydroxyvitamin D (D3) has been increased to 4000 IU/d, but levels of 10,000 IU/d for treatment of inflammatory conditions and 50,000 IU/week for short-term supplementation in deficient individuals have been suggested. High doses of D3 may cause arterial calcification and cardiomegaly; however, the blood velocity of the adverse events and related to regular D3 intake in doses higher than 10,000 IU/d, threshold for these changes and links to arterial calcification or hypertension are unclear. The objective of this study was to examine the effect of dietary doses of D3 equivalent to recommended human doses after 1 month in male rats (n = 4/group): 0 IU/d (deficiency), 600 (recommended daily intake; control), 10,000 (upper therapeutic dose), 30,000 & 150,000 IU/d on cardiovascular function. Blood pressure telemetry and carotid artery Doppler ultrasonography were done at the baseline and weekly during diet testing. After 4 weeks, both systolic and diastolic blood pressure increased significantly (~15 mmHg, ANOVA, Duncan’s Post-hoc test) in both D3 deficiency and groups receiving ≥30,000 IU/d compared to the 10,000 IU/d group. Carotid artery luminal diameter was significantly smaller in 0 or ≥30,000 IU/d D3 compared to control, which could indicate either increased arterial wall thickness or contractility. Pulse wave velocity and peak acceleration decreased over time in control and D3 deficiency, but increased in groups receiving ≥30,000 IU/d D3 indicating increased arterial stiffness. In conclusion, based on decreased blood pressure, caution needs to be taken with D3 supplementation ≥30,000 IU/d. Future studies will be directed at confirming the dose threshold and examining mechanisms for D3-induced arterial stiffness and hypertension.

**1128 Fatty Acids Down Regulate and Inhibit Secretion of Adiponectin in Adipocytes through Oxidative Stress.**

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Adipocyte tissue is not merely a storage depot for the fat, but has emerged as a vital organ capable of synthesizing and releasing adipokines which act as regulators of diverse physiological functions. Adiponectin (30 kDa) protein, an important member of the adipokine family is known to regulate insulin action, NADPH oxidase activity, and lipid mobilization. Adipose, release of free fatty acids from the adipose fat, and adiponectin secretion by the adipocyte are interconnected. These phenomena appear as crucial factors in cardiovascular and sleep disorders. Reports have been made on decreased circulating adiponectin levels in obese individuals, suggesting a link between fat load in adipose tissue and decrease in adiponectin release by adipose tissue. However, polyunsaturated fatty acids (PUFA) have been established as essential dietary fats and recently n-3 fatty acids (Omega-3) have gained importance in health, nutrition and disease. Having this as the premise, here we hypothesized that PUFA load would modulate adiponectin secretion by adipocytes. To test our hypothesis, we loaded the 3T3 differentiated adipocytes with different PUFAs including linoleic (C18:2), linolenic (C18:3), arachidonic (C20:4), eicosapentaenoic (20:5); EPA, and docosahexaenoic (22:6); DHA) acids for 24 h and then analyzed the secreted adiponectin, intracellular adiponectin, fatty acid composition, triglyceride accumulation, cytotoxicity, and extent of lipid peroxidation (formation of 4-hydroxy-2-nonenal, 4-HNE) in the cells. Our results revealed that PUFA loading suppressed both adiponectin release by the cells and intracellular adiponectin, increased cellular PUFA content and triglyceride accumulation, and induced lipid peroxidation (intracellular formation of 4-HNE). Overall, our results suggested that the essential PUFAs suppressed secretion and synthesis of adiponectin in adipocytes through elevated lipid peroxidation and oxidative stress.

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The regulatory guideline S7A supports the use of unrestrained models for safety pharmacology assays. Parenteral bolus administrations including the intravenous (IV) and subcutaneous (SC) routes are associated with CMAX achieved immediately or within minutes of dosing for some compounds. In this context, animal handling can constitute a major interference to data quality and can obscure pharmacodynamic responses. The current investigation compared cardiovascular changes after remote SC and IV delivery in telemetered cynomolgus monkeys and Beagle dogs with restrained administration in the same species. Remote parenteral administrations were associated with a substantial decrease in data variance and consequently improved minimum detectable differences. Mean heart rate variations from baseline in the first 30 min after restrained IV saline dosing reached a maximum of 27% compared to 9.7% for remote administration. Baseline systolic arterial pressure following remote SC saline dosing in cynomolgus monkeys ranged from 93.7 and 116.4 mmHg while post-dose values in the initial 60 min post-dose ranged from 94.0 and 116.3 mmHg. Recovery of cardiovascular parameters to baseline levels after animal handling was 25 to 120 minutes in both species which can constitute a major confounder to evaluation of short half-life drug candidates. Remote dosing presented optimal results when the animals remained completely undisturbed (i.e., no staff present in the animal room). Remote IV dosing is generally recognized to improve telemetry data quality but the current investigation also demonstrate beneficial effects of remote SC injections on sensitivity of cardiovascular investigations in safety pharmacology studies using telemetry in cynomolgus monkeys and Beagle dogs.

1131 Acute Silver Nanoparticle Exposure Increases Cardiac Ischemic/Reperfusion Injury in Sprague-Dawley Rats.

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The expanding use and production of silver nanoparticles (AgNP) as anti-bacterial/fungal agents is raising concerns regarding their safety to human health. The diversity of nanosized silver (AgNP) and coatings for dispersion may produce combinations that minimize potential toxic effects on the cardiovascular system. We hypothesized that acute intratracheal (IT) exposure to 20 nm (S) or 110 nm (L) AgNP coated with either polyvinylpyrrolidone (P) or citrate (C) would increase the susceptibility of cardiac tissue to a regional ischemic reperfusion (I/R) injury. Young male Sprague-Dawley rats were exposed to 200 µg of AgNP 24 hrs post-exposure, cardiac ischemia was induced for 20 mins followed by 2 hrs of reperfusion in situ. Hearts were sectioned stained with Evans blue to demarcate Area at Risk (AAR) and counter stained with TTC to determine % of AAR infarcted. Bronchoalveolar lavage fluid (BALF) and serum was collected post I/R injury to evaluate pulmonary injury and circulating markers of injury. Neither P (22±2% Infarct/AAR) nor C (24±1% Infarct/AAR) altered the extent of infarcted cardiac tissue as compared to naive (22±2% Infarct/AAR). However, the installation of all forms of AgNP significantly increased the extent of cardiac I/R injury. As a group the P-coated AgNP developed larger infarcts than C-coated. The 20 nm were more effective at enhancing I/R Injury (SP 43±2.0% Infarct/AAR; SC 37±3% Infarct/AAR) than the 110 nm size (LP 37±3% Infarct/AAR; LC 31±2% Infarct/AAR). The opposite pattern was observed in the BALF endpoints with P and L AgNP having greater effects. Our results suggest IT exposure to AgNP have differential effects on pulmonary and cardiac tissues following the application of I/R injury. This work is supported by NIEHS U19 ES019525.

1132 Effect of Pulmonary Exposure to Welding Fumes on Cardiomyocyte Contractility.

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Studies have found that pulmonary exposure to welding fumes is positively associated with a higher incidence of cardiovascular events. We reported previously that pulmonary exposure to welding fumes, manual metal arc-hard surfacing (MMA-HS), has a negative impact on cardiac function as evidenced by reduced heart contractility. However, the mechanisms underlying MMA-HS-induced depression of cardiac contractility remain unclear. To study the mechanisms, rats were given an intratracheal instillation of MMA-HS welding fumes (2 mg/rat) or saline once a week for seven weeks. Cardiomyocytes were isolated at 1 and 7 days post-exposure. Cardiomyocyte contractility and intracellular calcium level in response to increasing concentrations of adrenergic agonist isoprenaline and extracellular calcium were assessed using a Myocyte Calcium Imaging/Cell Length System. Pulmonary exposure to MMA-HS blunted contractile function in response to both isoprenaline and calcium at 1 day post-exposure (P<0.01; P<0.05, respectively). A blunted contractile response was also observed with welding fume treated rats at 7 days post-exposure in response to isoprenaline and calcium (P<0.01). Intracellular calcium level in response to extracellular calcium stimulation was reduced at 7 days post-exposure (P<0.05). These findings suggest that pulmonary exposure to welding fumes impairs cardiac function by decreasing cardiomyocyte contractility through a defect in the adrenergic signaling pathway and intracellular calcium handling.

1133 Silver Nanoparticle Exposure Increases Vasoconstrictor Response in Nonpregnant Female Sprague-Dawley Rats.

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The use and production of silver nanoparticles (AgNP) is growing rapidly raising concerns regarding their safety to human health, particularly following translocation to the circulatory system. Previous findings from our lab have shown intravenous (IV) exposure to AgNP changes the vasoconstrictor response in both pregnant and male Sprague-Dawley (SD) rats. We hypothesized, acute IV exposure to AgNP in non-pregnant females will increase vascular reactivity of arterial vessels.
from mesenteric and uterine vascular beds and aortas and these changes will be influenced by NP size and coating. Female Sprague-Dawley rats, 6 months old, were intratracheally exposed to 200 μg of 20 or 110 nm AgNP coated with polyvinylpyrrolidone (PVP) or citrate and suspended in water. 24 hrs. post-exposure, wire myography tested the vasomotor responses in aortic, first order mesenteric and main uterine artery segments. Cumulative dose response curves were created for phenylephrine (PE), angiotensin II (ANGII), and endothelin 1 (ET1). Segments of uterine artery and aorta from AgNP exposed animals generated larger stress when compared to vehicle controls. No significant differences were observed in the mesenteric artery responses. Maximum stress values in the uterine artery were greater (p < 0.05) in response to ET1 and ANGII stimulation following exposure to 110 nm PVP AgNP (16.4 ± 2.7 and 14.5 ± 2.2 mN/mm2, respectively) as compared to 110 nm citrate AgNP (8.0 ± 1.0 and 6.9 ± 0.8 mN/mm2, respectively). Conversely, maximum stress in response to PE in aortic segments was greater (p < 0.01) following exposure to 20 nm PVP AgNP (1.3 ± 0.2 mN/mm2) as compared to 20 citrate AgNP (2.8 ± 0.3 mN/mm2). Our results suggest IV exposure to AgNP has differential effects on increasing the vasocostructor responses that are dependent on the vascular bed, size of the NP and type of coating. This work is supported by NIEHS U19 ES019525.

1134 Intratracheal Administration of C60 Differentially Promotes Constriction or Impairs Relaxation of the Isolated Coronary Artery.

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The potential uses of C60 fullerene have grown to include roles in both commercial industry and medicine, but the impacts on human health are not completely understood. Data from our lab suggests that C60 can exacerbate cardiac ischemia/reperfusion injury. We hypothesized that exposure to C60 would promote enhanced vasoconstriction and impaired endothelium-dependent relaxation responses of the coronary artery. Male Sprague-Dawley rats were exposed to a single 93.3 μg/kg dose of intravenous (IV) or intratracheal (IT) C60 or polyvinylpyrrolidone (PVP) control. Twenty-four hours following exposures, coronary artery segments were isolated and evaluated using wire myography. Cumulative dose responses to serotonin (5-HT), acetylcholine (ACh), or sodium nitroprusside (SNP) were constructed. We found that IT exposure to C60 resulted in a leftward shift (P = 0.05) of the 5-HT EC50 (558.4 ± 104.5 nM) compared to vehicle (870.7 ± 129.7 nM), but that IV exposure to C60 did not significantly shift the 5-HT EC50 values (C60: 762 ± 112.3 μM vs. vehicle: 669.3 ± 59.4 μM). Conversely, IV exposure to C60 led to a rightward shift (P = 0.09) in the EC50 for ACh (232.5 ± 68.9 μM) compared to vehicle (100.9 ± 48.8 μM), but IT C60 exposure did not produce any differences in ACh EC50 (C60: 245.7 ± 37.4 μM compared to vehicle: 222.4 ± 31.6 μM). The relaxation responses to SNP were not different between any treatment groups. The route of exposure to C60 appears to influence an enhancement of smooth muscle contraction by IT and impaired endothelial relaxation by IV administration. Based on these data we conclude that C60 exposure could enhance coronary vascular tone, possibly compromising dilatory flow during reperfusion, and thus exacerbating myocardial infarction. This work is supported by NIEHS U19 ES019525.

1135 Intratracheal Exposure to Silver Nanoparticles Promotes Enhanced Coronary Vascular Tone.

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Silver nanoparticles (AgNP) have physicochemical and antimicrobial properties that make them useful in both industrial and biomedical applications. In regard to human exposures, little is known about the way AgNP may impact physiological systems. We tested the hypothesis that coronary responsiveness to serotonin (5-HT) and acetylcholine (ACh) would be augmented 24 hrs following exposure to AgNP, and be dependent on particle sizes and the dispersants used to coat the particles. We used male Sprague-Dawley rats (10-12 weeks old) and delivered AgNP (1 mg/kg), of either 20 nm (SS) or 110 nm (LS) diameters, and suspended in either polyvinylpyrrolidone (PVP) or citrate (Cit), by intratracheal (IT) instillation or intravenous (IV) injection. After 24 hrs we isolated segments of the left anterior descending coronary artery for wire myography analysis of 5-HT and ACh dose responses. Our data indicate no differences in coronary responses from AgNP groups compared to their vehicle groups. We did observe some unique dispentant- and AgNP size-dependent alterations in the coronary responses. Maximal 5-HT stresses were higher (P < 0.05) in the SSS-IT group (4.3 ± 0.5 mN/mm2) compared to the SSS-PVP-IT group (2.9 ± 0.3 mN/mm2) or the LSCit-IT group (3.2 ± 0.4 mN/mm2). ACh responses were impacted in a dispersant-dependent manner, shifting the EC50 from 106.6 ± 16.6 mN in the LSPVP-IT group to 50.4 ± 19.9 mN in the LSCit-IT group. We found no significant differences between coronary responses from rats exposed intravenously to AgNP or their vehicles. We conclude that pulmonary exposure to AgNP activates biological response pathways that are sensitive both the dispentant and AgNP size. This work is supported by NIH U19 ES019525.

1136 Environmental Predictors of Oxidized Ldl Cholesterol (oxLDL) in Navajo Populations Exposed to Uranium-Contaminated Mining Sites.

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Numerous abandoned mines within the Navajo Nation contribute uranium, arsenic and other heavy metals to the soil and groundwater. Environmental exposure to heavy metal contaminants may promote or exacerbate cardiovascular disease. The prevalence of type 2 diabetes, a risk factor for cardiovascular disease, has increased among the Navajo community. Evidence is emerging that pre-existing metabolic disease may increase the risk of indices of toxicity of heavy metal contaminants. To assess the potential impact of these contaminants on cardiovascular health of exposed individuals, we examined traditional (IL-6, CRP) and novel (oxLDL, LOX-1 receptor) plasma biomarkers in a large community of the Navajo Nation. Samples and data were obtained through a culturally appropriate community-based participatory approach, incorporating data collection and outreach by Navajo community staff. Biomarker data were linked to geospatial data on contamination sites using traditional linear regression and Bayesian models. When we used a binary model for uranium and arsenic drinking water levels, we observed that the estimated annual intake of arsenic was a significant predictor of oxLDL; age, the occupational exposure score, and the distance-environmental exposure score were also significant predictors. Diabetes, based on glycated hemoglobin, was a significant predictor of oxLDL. No environmental factors were correlated with LOX-1, CRP or IL6. In summary, oxLDL does not seem to trend with environmental exposure to abandoned uranium mines or their heavy metal contaminants, but oxLDL does seem to trend with arsenic in drinking water. While still preliminary, these results indicate that arsenic intake may increase markers of cardiovascular risk.

1137 Pulmonary Nanoparticle Exposure Enhances Cardiac Parasympathetic Signaling.

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Pulmonary nanoparticle exposure has been associated with alterations in autonomic signaling in the heart following particle exposure. We have recently reported that nanoparticle exposure alters microvascular responses to sympathetic nerve stimuli. Thus we developed a model of baroreceptor function reflexes to determine the relative autonomic contributions. Rats were exposed to titanium dioxide (nano-TiO2) for 4 hr for 1 day at 9 mg/m3, 24-hours post-exposure, baroreflex sensitivity was determined by i.v. infusion of sodium nitroprusside (SNP; 5-20 μg/kg) or phenylephrine (PE; 1-16 μg/kg), with or without atropine (1 mg/kg) to diminish parasympathetic signaling. Nano-TiO2 exposure did not alter PE-induced (4 μg/kg) changes in mean arterial pressure (AMAP; 25±5 mm Hg Sham, 26±3 mm Hg Exposed). However, nano-TiO2 exposure increased the magnitude of Ahearte rate in beats per minute (AMAP; 20±6 control vs. 26±5 exposed), suggesting an enhanced parasympathetic response. When these experiments were repeated in the presence of atropine, PE-induced ABPM was diminished in both groups; however atropine significantly altered ABPM in nano-TiO2, compared to sham (-18±1 Sham, -12±2 Exposed), suggesting an enhanced sensitivity to muscarinic receptor blockade. Atropine treatment did not significantly alter AMAP in either group. Nano-TiO2, exposure did not significantly alter SNP-induced changes in AMAP (-36±5 μM Hg control vs. -39±5 μM Exposed), or heart rate (ABPM 18±2 control vs. 26±7 exposed). Because parasympathetic projections in cardiovascular system are largely confined to the heart, these data suggest that nano-TiO2 exposure
1138 Cigarette Smoke Induces Ventricular Remodeling through Activation of the Aryl Hydrocarbon Receptor.

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Cigarette smoking is the major cause of preventable morbidity and mortality, with 5 million deaths annually worldwide. Cigarette smoke (CS) increases the risk of cardiovascular disease, including myocardial infarction and coronary artery disease. Of the 4700 identified components in CS, 60 are known carcinogens with polycyclic aromatic hydrocarbons (PAH) the most abundant carcinogenic agent. Chronic exposure to PAH can lead to ventricular dilation and dysfunction. Previously, we found the 5 wks of exposure to diesel exhaust particulates (DEP), which contain a high level of PAH similar to that in CS, induced ventricular dilation and dysfunction. DEP exposure impaired ventricular remodeling through activation of the aryl hydrocarbon receptor (AHR). Because of the similarities in chemical composition, we hypothesized that chronic exposure to CS would activate the AHR pathway inducing ventricular extracellular matrix (ECM) remodeling. Male Sprague Dawley rats were exposed to six cigarettes per day (Kentucky 2R4F) for 12 wks using a modified version of the Griffith snout exposure method. CS exposure resulted in increased expression of AHR in the left ventricle. Furthermore, these animals had increased cardiac expression of cytochrome P450 1A1, an enzyme upregulated by AHR activation. CS exposure reduced cardiac collagen as assessed through decreased levels of hydroxyproline. In addition, CS increased the expression of MMP-9, a key regulator in collagen turnover. These in vivo findings were confirmed in isolated cardiac fibroblasts, where 10% CS extract reduced the secretion of collagen by cardiac fibroblasts. In all, our findings indicate that CS alters ventricular ECM remodeling through the reduction of myocardial collagen by an AHR dependent mechanism.

1139 The Association of Vascular Disease with Exposure to Diesel Exhaust.

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Cardiovascular disease (CVD) includes numerous heart- and circulatory-system-related conditions, many of which are related to the buildup of plaque in the walls of coronary (heart) arteries, in which case the disease is coronary artery disease (CAD). CVD is the leading cause of mortality in the US, with about 25% of all deaths related to some form of this disease. While the majority of research on the health effects of exposure to diease (DE) has focused on pulmonary end-points, there is a growing body of research on the cardiovascular effects of DE particulate inhalation. Because DE exposure in the US is widespread, both occupationally and regionally, possible cardiovascular effects from DE could have significant public health implications. We performed a weight-of-evidence evaluation of the potential for DE to cause CAD, considering the mechanistic, toxicological, and epidemiological aspects of DE. Based on our research and analysis, we find little evidence to substantiate an association of DE exposure with CAD (ambient) exposure levels. In particular, the occupational epidemiology studies fail to provide consistent evidence of increased risk. We identified twelve studies of diesel-exposed worker populations in which cardiovascular health endpoints were studied. Several of these studies found significantly increased risks of one or two endpoints (or in one or two subpopulations), while other studies reported significantly decreased risks for some endpoints/subpopulations. Overall, there was not a consistent effect observed across the literature. Moreover, because of a lack of dose-response between DE particulate levels and CVD mortality, the results of our evaluation also have implications for assigning cardiovascular mortality risk to ambient air PM2.5, as is often done in assessing monetary value of PM2.5 reductions.

1140 FGF21 Expresses in Diabetic Hearts and Protects from Palmitate- and Diabetes-Induced Cardiac Cell Death In Vitro and In Vivo via Erk1/2-Dependent p38 MAPK/AMPK Signaling Pathway.

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The present study examined whether fibroblast growth factor (FGF) 21 expresses in the heart of diabetic mice and also protects from fatty acid (palmitate)- and diabetes-induced cardiac apoptosis. Stretboxnotin (STZ) induced type 1 diabetes increased FGF21 expression about 40 fold at 2 months and 1.5 fold at 6 months. To define if the up-regulated cardiac FGF21 expression offers a protective effect on fatty acid- or diabetes-induced cardiac damage, H9C2 cells were exposed to palmitate at 62.5 μM for 15 h, which significantly increased apoptosis. Pre-incubation of palmitate-treated cells with FGF21 significantly reduced the apoptosis, examined by DNA fragmentation and cleaved caspase-3. Mechanistically FGF21 inhibited palmitate down-regulated phosphorylation levels of Erk1/2, p38 MAPK and AMPK. Via application of inhibitor of each kinase and Erk1/2 sRNA, FGF21 was found to prevent palmitate-induced cardiac apoptosis via up-regulating Erk1/2 dependent p38 MAPK/AMPK signaling pathway. To confirm these in vivo, STZ-induced diabetic mice were treated with FGF21 for 10 days, which significantly prevented diabetes-induced cardiac apoptosis along with up-regulation of Erk1/2, p38 MAPK and AMPK phosphorylation. The cardiac protection of FGF21 was completely blocked by Erk1/2 inhibitor. FGF21 also protected cardiac apoptosis in acute fatty acid-treated mice via the same signaling pathway. More importantly, compared to wild-type mice, FGF21 gene knock-out mice are highly susceptible to diabetes-induced cardiac cell death and inhibition of Erk1/2, p38 MAPK and AMPK phosphorylation, which could be prevented by FGF21 treatment. These results strongly suggest that the early stage increase in FGF21 mRNA expression may represent an early protective mechanism: Application of FGF21 to diabetic mice or acute fatty acid-treated mice can prevent cardiac cell death via Erk1/2-dependent p38 MAPK and AMPK signaling pathway.

1141 Effects of Nicotine on Cardiovascular Remodeling in a Mouse Model of Systemic Hypertension.

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The association of cigarette smoking and hypertension with the development of heart disease and aortic aneurysms is clear. The use of nicotine-alone formulations, such as the nicotine patch, gum or ‘smokeless’ electric cigarettes is increasing, as they are perceived as healthier alternatives to traditional cigarettes. Unfortunately, there is little data available on the effect of isolated nicotine on myocardial and aortic remodeling in healthy subjects or in the setting of hypertension. We hypothesized that nicotine would exacerbate the effects angiotensin-induced hypertension, as evidenced by reduced left ventricular wall thickness and aortic wall remodeling. For this study we utilized subcutaneous osmotic minipumps to administer angiotensin II, nicotine, nicotine plus angiotensin II (Ang-II) or saline to C57Bl/6 (n=6) mice for a total of 4 wks. Ventricular wall thickness, activity of matrix metalloprotease 2 and 9 (MMP2/9) and presence of type 1 collagen was assessed as evidence of cardiac remodeling. Heart weights were increased by treatments, with control<nicotine<ang-II<nicotine+ang-II, but a statistical interaction of nicotine x ang-II was not noted. Activity levels of MMP-2 mirrored these changes and demonstrated a calculated additive effect of nicotine and ang-II. Additionally, histopathological analysis of aortas revealed that combined nicotine and ang-II treatment induced significant hypertrophy compared to all other groups. This pilot project reveals possible cardio toxic interactions between nicotine and angiotensin II hypertension. These data indicate that in the presence of hypertension, nicotine alone can drive development of cardiovascular disease, which has important implications on labeling and marketing of nicotine replacement therapy. These data also raise the possibility that angiotensin inhibitors may be specifically efficacious in prevention of cardiovascular disease among hypertensive individuals using nicotine.

1142 Metallothionein Preservation of Akt2 Function by Down-regulating TRB3 Restores Diabetic Inhibition of Cardiac Insulin Signaling.

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Cardiac insulin resistance is a key pathogenic factor for diabetic cardiomyopathy, but its mechanism is largely unclear. Here we demonstrated that streptozocin-induced diabetes significantly inhibited protein kinase B (Akt) Ser473 and Thr308 phosphorylation from 2 weeks to 2 months. In cardiac-specific metallothionein-transgenic (MT-TG) diabetic mice, both Akt phosphorylation sites were down-regulated with marked inhibition of Akt phosphorylation. Akt phosphorylation can be increased through the Wnt/β-catenin pathway, while Akt phosphorylation can be inhibited by the tribbles (TRB3) expression. The present study indicates that the MT-TG model may be a useful tool to study the mechanism of Akt phosphorylation and cardiac insulin resistance.

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Acute drug-induced vascular injury (DIVI) is caused by several classes of drugs, including Phosphodiesterase (PDE) inhibitors and dopamine agonists. The finding of DIVI in preclinical toxicity studies presents a significant challenge for pharmaceutical companies, with many compounds terminated from development because of DIVI in preclinical toxicity studies. Exposure to fine particles (PM2.5) is associated with adverse cardiopulmonary health effects; however, the causative components in PM2.5 are limited. Nickel is one of the major elements of ambient particulate matters. This study was designed to investigate the acute cardiovascular systemic toxicity induced by PM2.5 on Wistar rats and the role of nickel sulfate in it. Male Wistar rats were exposed by intratracheal instillation to blank membrane (control), fine particles with the doses of 7.5, 15 and 30μg/kg body weight and balanced saline (control), nickel sulfate dosages of 7.5, 75 and 750μg/kg Ni; The rats were sacrificed after 24h, blood samples were collected and parameters of inflammation, cytokines, coagulative effects and the nickel levels in serum were measured to estimate the cardiovascular injury. The results showed that both of PM2.5 and Nickel sulfateinduced a significant increase of serum C-reactive protein, IL-6 and TNF-tt, PT levels in the plasma decreased significantly in PM2.5-treated rats compared with the control group. Plasma fibrinogen increased significantly at 15 μg/kg and 30 μg/kg PM2.5 groups. The PT and APTT showed no changes in NISO4-treated rats, Plasma fibrinogen levels increased significantly at 750 μgNi/kg NISO4-treated rats. The TF concentrations in plasma increased significantly at 75 μgNi/kg and 750 μgNi/kg dosage groups (P<0.05), there was a slight increase at low dosage (7.5 μgNi/kg) group. To be compared with the corresponding concentration of NISO4, the acute effects of PM2.5 were more pronounced. Our results suggest that water-soluble nickel in PM2.5 may play an important role in PM2.5 toxic effects on animal cardiovascular system. Key words: PM2.5; Nickel sulfate ; CardiovascularToxicity

Note: This experiment was carried out in accordance with the Peking University Health Sciences Center, New Orleans, LA.

1143 Development of an Ultrasound Imaging Biomarker of Drug-Induced Vascular Injury.

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Acute drug-induced vascular injury (DIVI) is caused by several classes of drugs, including Phosphodiesterase (PDE) inhibitors and dopamine agonists. The finding of DIVI in preclinical toxicity studies presents a significant challenge for pharmaceutical companies, with many compounds terminated from development because of the inability to monitor DIVI in the clinic. Ultrasound imaging is a non-invasive technique that can be used to measure blood flow and vessel diameter in multiple tissues and across species, and therefore has the potential to be a translatable biomarker of DIVI. Our objective was to demonstrate the utility of high-frequency ultrasound imaging for measuring hemodynamic changes in mesenteric arteries of male Sprague Dawley rats treated with fenoldopam and CI-1044 (PDE4 inhibitor). Blood flow and vessel diameter were measured in the superior mesenteric arteries and right renal arteries at 4, 8 and 24 hours after dosing and the rats were subsequently necropsied at each time point. Microscopic observations consistent with DIVI were found in animals treated with fenoldopam at 24 hours and included mild to moderate perivascular accumulations of mononuclear cells, neutrophils in tunica adventitia and superficial tunica media as well as multifocal hemorrhage and necrosis in the tunica media. Animals dosed with both drugs showed marked increases in flow and shear stress as early as 4 hours after dosing, with the fenoldopam-treated rats exhibiting the largest changes in these parameters. Vessel diameter and blood flow were used to calculate shear stress and regression analysis was used to model the relationship between the presence of lesions and various hemodynamic parameters and indicated that peak blood flow velocity was the best predictor of lesion formation. This data suggests that ultrasound imaging may provide a translatable and non-invasive functional biomarker for localized hemodynamic changes that may correlate with acute vascular lesion formation in specific target tissues.

1144 Environmentally-Persistent Free Radicals (EPFRs) Exacerbate Cardiac Dysfunction during Ischemia-Reperfusion (I-R) Injury.

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EPFRs have been identified in the particulate matter (PM) milieu at Superfund sites. These EPFRs, capable of redox cycling and the continual generation of radical species, form during combustion processes as halogenated hydrocarbons chemisorb to transition metal oxide-containing particles. We showed that short-term nose-only inhalation of EPFRs decreased stroke volume (SV) and cardiac output (CO) in rats secondary to increased pulmonary resistance and decreased ventricular filling. EPFRs also increased markers of oxidative stress and inflammation both systemically and in the left ventricle. Epidemiological studies link PM exposure to increased cardiac morbidity and mortality from ischemic events. Therefore, the current study sought to determine if prior exposure to EPFRs would increase cardiac vulnerability to I-R injury. The EPFRs (0.2 μm diameter) were surrogates consisting of 1,2-dichlorobenzene chemisorbed to a silica/CoO particle at 230°C (DCB230). Rats were exposed via nose-only inhalation to DCB230 (171 μg), silica particles or vehicle for 20 min/day for 7 days. Ventricular function was measured in vivo using pressure-volume catheters. I-R injury was induced by ligating the left anterior descending coronary artery for 20 min followed by 60 min of reperfusion.

Compared to controls, baseline CO, stroke work and SV were significantly reduced by exposure to DCB230. These parameters were further reduced during ischemia and remained below control throughout reperfusion. Although not significantly different from controls, end diastolic volume and heart rate trended lower in DCB230 treated rats throughout, possibly contributing to the reduced ventricular function. Indices of contractility and diastolic function were not different between the groups. These data show that EPFRs reduce baseline ventricular function and exacerbate the decrease in ventricular function elicited during I-R injury. Supported by NIH P42-ES013648 (subaward 61365).

1145 The Role of Nickel in Systemic Inflammation and Blood Coagulative Effects Induced by PM2.5.

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Exposure to fine particles (PM2.5) is associated with adverse cardiopulmonary health effects; however, the causative components in PM2.5 are limited. Nickel is one of the major elements of ambient particulate matters. This study was designed to investigate the acute cardiovascular systemic toxicity induced by PM2.5 on Wistar rats and the role of nickel sulfate in it. Male Wistar rats were exposed by intratracheal instillation to blank membrane (control), fine particles with the doses of 7.5, 15 and 30μg/kg body weight and balanced saline (control), nickel sulfate dosages of 7.5, 75 and 750μg/kg Ni; The rats were sacrificed after 24h, blood samples were collected and parameters of inflammation, cytokines, coagulative effects and the nickel levels in serum were measured to estimate the cardiovascular injury. The results showed that both of PM2.5 and Nickel sulfateinduced a significant increase of serum C-reactive protein, IL-6 and TNF-tt, PT levels in the plasma decreased significantly in PM2.5-treated rats compared with the control group. Plasma fibrinogen increased significantly at 15 μg/kg and 30 μg/kg PM2.5 groups. The PT and APTT showed no changes in NISO4-treated rats, Plasma fibrinogen levels increased significantly at 750 μgNi/kg NISO4-treated rats. The TF concentrations in plasma increased significantly at 75 μgNi/kg and 750 μgNi/kg dosage groups (P<0.05), there was a slight increase at low dosage (7.5 μgNi/kg) group. To be compared with the corresponding concentration of NISO4, the acute effects of PM2.5 were more pronounced. Our results suggest that water-soluble nickel in PM2.5 may play an important role in PM2.5 toxic effects on animal cardiovascular system. Key words: PM2.5; Nickel sulfate ; CardiovascularToxicity

Note: This experiment was carried out in accordance with the Peking University Health Sciences Center, New Orleans, LA.

1146 Involvement of Shear Stress in Fenoldopam and Dopamine-Induced Mesenteric Medial Arterial Necrosis.


Extrapulmonary and relevance of drug-induced rat vascular injury to humans has hampered drug development due to lack of understanding of the pathologic mechanisms involved. Although vasodilatation and increased shear stress (SS) have been hypothesized to be involved in the pathogenesis of these lesions, the exact role of SS on primary target cells, vascular smooth muscle (VSMC) and endothelial cells (EC) in vivo remain unclear. Dopaminergic D1 agonists such as Fenoldopam and Dopamine reproducibly induce mesenteric medial arterial necrosis (MAN) in rats following single vasotoxic doses. To investigate the involvement of SS in the development of MAN, rats were given vehicle, Dopamine or Fenoldopam for 4 days. Yohimbine, a alpha-2 adrenoreceptor antagonist, also vasoactive but lacking MAN was given to rats for 4 days for comparison. To evaluate the timecourse of lesion development, rats were also given vehicle or a single vasotoxic dose of Fenoldopam and necropsied 1-, 4-, 6-, 12- or 24 hours postdose. Mesentry from each rat was collected and frozen in OCT, then EC and VSMC were microdissected from rat mesenteric arteries, and RNA was amplified and analyzed. Regulations of 37 shear stress responsive genes were evaluated using TaqMan® gene expression profiling. Many of the genes evaluated were confirmed to be differentially expressed by Dopamine and Fenoldopam in EC- and/or SMC-enriched samples as compared to controls and Yohimbine following 4 daily doses. A number of SS responsive genes were also shown to be differentially regulated beginning 1- and/or 4-hours post-Fenoldopam treatment and prior to histological evidence of MAN (which was initially observed beginning at 12-hours). Evaluation of this panel of genes has provided evidence of the involvement and regulation of SS responsive genes in both EC and VSMC during the development of Dopamine and Fenoldopam-induced vascular injury.
Direct Interaction between Multiwalled Carbon Nanotubes and the Coronary Microcirculation.

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Recently our collaborative group identified multi-walled carbon nanotube (MWCNT) translocation from the lung to the heart, liver, and kidneys within 24 hours after pulmonary exposure. From this finding, we continued to examine the microvascular ramifications associated with this direct interaction. To model drug delivery platforms, as well as lung migration, MWCNT were injected into the tail vein of rats (25-900 μg, suspended in 900 μL normalol). 24-hours later, coronary arteries (<170 μm in diameter) from the left anterior descending artery distribution were isolated for reactivity assessments based on responses to transmural pressure (myogenic responsiveness), intraluminal flow (shear stress), phenylephrine (10-9-10-4 M), acetycholine (10-9-10-4 M), A23187 (10-9-10-5 M), and spermineNONOate (10-9-10-4 M). Myogenic responsiveness was not altered after MWCNT injection. However, MWCNT injection at all concentrations robustly attenuated reactivity. The coronary microvascular dysfunction associated with MWCNT injection is significant, impacting endothelium-dependent, -independent, adrenergic, and flow-mediated dilation pathways. These alterations, in combination with previous findings, indicate that the microvascular impairments that follow MWCNT injection are more severe than those observed after ingestion or inhalation exposure. Studies are currently underway to further evaluate mechanistic differences between these routes of exposure.

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Phosgene-Induced Lethal Lung Edema Correlate with Persistent Stimulation of Cardiopulmonary Reflexes.

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Phosgene gas is a lower respiratory tract irritant. As such, it stimulates nociceptive vagal C-fiber related reflexes in a dose-rate and concentration x exposure duration (Cxt)-dependent manner. In rats this reflex is characterized by extended apnea time periods, bradycardia, and hypothermia. While inhalation exposures at non-lethal Cxt products show rapid reversibility of reflexively induced changes in respiratory patterns, lethal Cxt products seem to cause their prolonged stimulation after discontinuation exposure to phosgene. This observation has been taken as indirect evidence that phosgene-induced lethal lung edema is likely to be caused by dysfunctional neurogenic control of cardiopulmonary and microvascular physiology. In order to verify this hypothesis, data from respiratory function measurements during and after the inhalation exposure to phosgene gas were compared with time-course measurements of respiratory and cardiac function over 24 hours post-exposure. These data were complemented by time-course analyses of nitric oxide (NOe) and carbon dioxide in exhaled breath, including time-dependent changes of NOe in exhaled breath following acute high-level exposure. These data were complemented by time-course analyses of nitric oxide and carbon dioxide in exhaled breath, including time-dependent changes of NOe in exhaled breath following acute high-level exposure. These data were complemented by time-course analyses of nitric oxide and carbon dioxide in exhaled breath, including time-dependent changes of NOe in exhaled breath following acute high-level exposure. These data were complemented by time-course analyses of nitric oxide and carbon dioxide in exhaled breath, including time-dependent changes of NOe in exhaled breath following acute high-level exposure.

Persistent Stimulation of Cardiopulmonary Reflexes.

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Exposure to trace metals can disrupt olfactory function in fish leading to a loss of behaviors critical to survival. Cadmium (Cd) is an olfactory toxicant that elicits cellular oxidative stress as a mechanism of toxicity while also inducing protective cellular antioxidant defenses. Translocation of the nuclear factor erythroid-derived 2-like 2 (Nrf2) pathway. However, the molecular mechanisms of Cd-induced olfactory injury have not been characterized. In the present study, we investigated the role of the Nrf2-mediated antioxidant defense pathway in protecting against Cd-induced olfactory injury in zebrafish. A dose-dependent induction of Nrf2 was observed. The inhibition of antioxidant gene induction in Cd-exposed Nrf2 morphants was associated with disruption of olfactory driven behaviors, increased cell death and loss of olfactory sensory neurons (OSNs). Nrf2 morphants also exhibited a downregulation of OSN-specific gene expression, suggesting a role for Nrf2 in regulating olfactory responses. Pre-treatment with sulforaphane (SPN) partially protected against Cd-induced olfactory tissue damage. Collectively, our results indicate that oxidative stress is an important mechanism of Cd-mediated injury in the zebrafish olfactory system. Moreover, the Nrf2 pathway plays a protective role against cellular oxidative damage and is important in maintaining zebrafish olfactory function. This work was supported by the University of Washington Supersfund Research Program (NIH P42ES004696).

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Role of Nrf2 Antioxidant Defense in Mitigating Cadmium-Induced Oxidative Stress in the Olfactory System of Zebrafish.

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Cadmium (Cd), known to be a causative agent of Ito-iitai disease, produces severe toxic effects in the kidney, liver, lung and bone. Particularly, chronic exposure to Cd causes renal dysfunction. Recently, we have found that overaccumulation of p53 may relate to Cd-induced apoptosis and may be due to the suppression of p53 degradation through the inhibition of gene expression of Ubc2d family in rat proximal tubular cells (NRK-52E cells). In this study, we examined the effect of Cd on the expressions of UBE2D family, accumulation of p53 and apoptosis in human proximal tubular cells (HK-2 cells). TUNEL positive cells, indicators of apoptosis, were increased by treatment with 20 μM Cd for 18 h. Moreover, using real-time RT-PCR, we demonstrated that Cd caused significant decrease of UBE2D2 mRNA levels from 6-h treatment, and UBE2D4 mRNA levels from 12-h treatment. Western blot analysis showed that Cd drastically increased p53 protein levels in HK-2 cells, even though the mRNA levels of p53 were decreased by Cd. Additionally, knockdown of UBE2D family genes by siRNA increased cellular protein levels of p53. These results indicate that Cd induces apoptosis through p53 accumulation by suppressions of UBE2D family gene expressions, in human proximal tubular cells as well as in rat proximal tubular cells.

PS 1147

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1150 Cadmium Enhances Instability of Lysyl Oxidase Messenger RNA.

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Lysyl oxidase (LO), a copper-dependent enzyme, catalyzes crosslinking of collagen and elastin essential for tissue and organ morphogenesis and repair. Our previous studies have shown the critical role of downregulation of LO gene in cadmium (Cd)-induced emphysema pathogenesis in vitro and in vivo. The present studies further investigate Cd effects on posttranscriptional modification of the LO gene in rat fetal lung fibroblasts (RFL6). Treatment of cells with Cd (1-5 μM) inhibited levels of LO mRNA stability as assessed by the actinomycin D (an inhibitor of mRNA synthesis, 5 μg/ml) chase assay with the t1/2 = 24 h. In contrast, in cells treated with 5 μM Cd plus actinomycin D, the t1/2 for LO mRNA decay was reduced to 0.75-1 h (45 min). Thus, Cd facilitates the LO mRNA decay. Rat LO mRNA contains two AU-rich elements (ARE, AUAUA), the stability determinant, at 174/178 and 200/204 downstream of the translation stop codon in the 3'-UTR. We cloned the entire LO 3'-UTR (1/229) and the ARE fragment (152/229) into the pGL3-Reporter vector after the coding region of the SV40 promoter-driven luciferase gene. The LO 3'-UTR or the ARE fragment strongly stabilized the reporter mRNAs manifested by increased luciferase activities in transfected cells. Mutation of two AREs abolished the enhancement of reporter mRNA stabilities by the LO 3'-UTR or the ARE fragment. Notably, luciferase activities driven by the LO 3'-UTR or ARE in recombinant constructs were significantly inhibited in cells treated with 5 μM Cd, indicating Cd enhancement of the decay of LO 3'-UTR or ARE-driven reporter mRNAs. HuR, a major ARE binding protein, enhances the mRNA stability. RNA immunoprecipitation (RIP) assay indicated that Cd effectively blocked the binding of HuR into the LO mRNA AREs. Thus, Cd targeting the HuR may be a critical mechanism for LO mRNA instability by this metal ion (supported by the grant of NIEHS 113140).

1151 Involvement of Inhibition of UBE2D Family Gene Expressions on Cadmium-Induced p53 Dependent Apoptosis in Human Proximal Tubular Cells.

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Cadmium (Cd), known to be a causative agent of Itai-itai disease, produces severe toxic effects in the kidney, liver, lung and bone. Particularly, chronic exposure to Cd causes renal dysfunction. Recently, we have found that overaccumulation of p53 may relate to Cd-induced apoptosis and may be due to the suppression of p53 degradation through the inhibition of gene expression of Ubc2d family in rat proximal tubular cells (NRK-52E cells). In this study, we examined the effects of Cd on the expressions of UBE2D family, accumulation of p53 and apoptosis in human proximal tubular cells (HK-2 cells). TUNEL positive cells, indicators of apoptosis, were increased by treatment with 20 μM Cd for 18 h. Moreover, using real-time RT-PCR, we demonstrated that Cd caused significant decrease of UBE2D2 mRNA levels from 6-h treatment, and UBE2D4 mRNA levels from 12-h treatment. Western blot analysis showed that Cd drastically increased p53 protein levels in HK-2 cells, even though the mRNA levels of p53 were decreased by Cd. Additionally, knockdown of UBE2D family genes by siRNA increased cellular protein levels of p53. These results indicate that Cd induces apoptosis through p53 accumulation by suppressions of UBE2D family gene expressions, in human proximal tubular cells as well as in rat proximal tubular cells.
Changes of serum and hepatic alkaline phosphatase activity in rats administered with 65 ppm of Cd2+ in drinking water were studied during a 3 months period. After metal administration, Cd2+ in rat livers showed an accumulation of 21 fold in 1 month Cd-exposed rats, however, later enzymatic activity decreased in the rats exposed to the metal by 3 months. In order to determine if Cd accumulated in liver had an inhibitory effect on enzyme activity, hepatic alkaline phosphatase from non-exposed and 3-month exposed rats to Cd was isolated and apo-alkaline phosphatase was reactivated with Zn2+ and Mg2+. Reactivation assays shown that liver alkaline phosphatase isolated from Cd-exposed rats had 2-fold more activity as compared with the enzyme isolated from Cd-non-exposed rats. Protein alkaline phosphatase did not show changes between non-exposed and metal-exposed rats as determined by western-blot and ELISA assays. Results of this work suggest that chronic Cd-exposure has an inhibitory effect on serum and hepatic alkaline phosphatase activity, and the mechanism of inhibition can involve the substitution of native zinc in the enzyme by the Cd progressively accumulated in the tissue.

Serum alkaline phosphatase activity decrease in 3 months Cd-exposed rats. Liver alkaline phosphatase activity increased after 1 month Cd-exposed rats, however, later enzymatic activity decreased in the rats exposed to the metal by 3 months. Liver alkaline phosphatase activity increased after 1 month Cd-exposed rats, however, later enzymatic activity decreased in the rats exposed to the metal by 3 months. In order to determine if Cd accumulated in liver had an inhibitory effect on enzyme activity, hepatic alkaline phosphatase from non-exposed and 3-month exposed rats to Cd was isolated and apo-alkaline phosphatase was reactivated with Zn2+ and Mg2+. Reactivation assays shown that liver alkaline phosphatase isolated from Cd-exposed rats had 2-fold more activity as compared with the enzyme isolated from Cd-non-exposed rats. Protein alkaline phosphatase did not show changes between non-exposed and metal-exposed rats as determined by western-blot and ELISA assays. Results of this work suggest that chronic Cd-exposure has an inhibitory effect on serum and hepatic alkaline phosphatase activity, and the mechanism of inhibition can involve the substitution of native zinc in the enzyme by the Cd progressively accumulated in the tissue.

Cadmium(Cd), a toxic metal initially accumulated in the liver, forms high affinity complexes with metallothionein-II (MT-II). This protein requires the activation of transcription factors such as STAT3. Phosphorylation is crucial to regulate the activity of several proteins, as well as acetylation, has been linked in gene expression processes giving greater affinity for the transcription factors to DNA. This process has been reported in STAT3 to form more stable dimers. There is evidence that this modification is carried out by the p300 coactivator protein, which is characteristic because of its histone acetyltransferase activity and its Stat3 C-terminal affinity. MT-II protein also has been associated with Src; however, molecular mechanisms induced by Cd have not been yet fully understood. The aim was to evaluate the participation of Src in Stat3 activation and its relation with MT-II production and acetylation in hepatocytes treated with 5μM CdCl2. Primary mouse hepatocytes were treated with different Cd concentrations determining phosphorylation of STAT3, Src, ERK1/2 and the induction of p300 and MT-II by Western-blot. A pretreatment for 30 min with Src inhibitor SU6656 was performed to evaluate the relationship between STAT3 activation and MT-II content; the same approach was done with the ERK1/2 inhibitor PD98059, used to correlate STAT3 activation and ERK signaling. We proved that Cd induced STAT3 acetylation. Cd increased activation of STAT3, Src, ERK1/2, p300, and MT-II STAT3 phosphorylation and ERK1/2 were significantly reduced by Src, ERK1/2 inhibitors. MT-II expression diminished with Src inhibitor. Our results suggest that Cd induces STAT3 phosphorylation by a mechanism dependent of Src and ERK1/2, while STAT3 induces acetylation by p300 likely to increase production of MT-II as a biological response to Cd.

Serum and Hepatic Alkaline Phosphatase Activity in Rats Exposed Chronically to Cadmium.

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Exposure to the carcinogenic metal cadmium can activate multiple signaling pathways to affect the expression of hundreds of genes. However, the molecular mechanism by which cadmium affects transcription is not completely understood. Whole genome microarray studies in the nematode Caenorhabditis elegans showed nuclear metal-responsive gene-1 (numr-1) mRNA levels increased 7-fold following a 24h cadmium exposure. Subsequent genomic analysis identified a second gene, numr-2, that is 99% identical to numr-1, in both coding and regulatory regions. Both numr-1 and numr-2 are developmentally regulated and expressed in identical cells: constitutively in a subset of neurons in the head, vulva and tail; and in intestinal and pharyngeal cells following metal exposure. Metals and calcium, but not other environmental stressors, induce numr-1/-2 transcription. Bioinformatic analysis identified 25 putative, upstream regulatory elements in the numr promoters. The roles of the cognate DNA binding proteins on numr-1 expression in the absence and presence of cadmium was examined in loss of function mutants or by RNAi. Among the tested genes, two alleles of omm-9 (TRPV channel protein), ek1677 and ky10, showed significant gene-cadmium interactions, suggesting omm-9 regulates cadmium-induced numr-1/-2 transcription. Three other genes may also be involved in numr-1/-2 transcriptional regulation: ceh-20 (homeodomain protein), unc-86 (POU-type homeodomain protein) and gem-4 (calcium-dependent phosphatidylinositol binding protein). In addition to omm-9, unc-86 and ceh-20 are expressed in neurons, suggesting a link between neuronal activity and metal-inducible numr-1/-2 transcription. The four candidate genes are present in independent signaling pathways, which suggest that cadmium-induced numr-1/-2 transcription is regulated by multiple signaling pathways. Further analyses disrupting these signal transduction pathways will be used to define the role of these genes in cadmium-induced numr-1/-2 transcription.
Diabetes mellitus (DM) and diabetes-related kidney disease are serious, worldwide health problems. Although there is no direct evidence linking cadmium (Cd) to DM, CD exposure alters blood glucose levels and potentiates diabetic nephropathy. The insulin/insulin-like growth factor signaling (IIS) pathway regulates multiple biological functions including glucose metabolism and longevity. The model organism C. elegans, which has an IIS pathway homologous to that of mammalian species, was used to investigate mechanistic links among Cd, transcription, and insulin signaling. The focus of this investigation was the C. elegans Cd-responsive gene cdr-1, whose transcription is up-regulated almost 800-fold exclusively by Cd. To identify regulatory factors and pathways that control cdr-1 transcription, an integrative transgenic strain of C. elegans containing GFP under the control of the 5' regulatory region of cdr-1 was constructed. Following cadmium exposure (10 μM, 24 h) in the transgenic strain, GFP expression increases in intestinal cells. In a candidate screen, genes involved in various stress response pathways were tested for their potential role in controlling cdr-1 expression. Genes were knocked out either by genetic crosses to known loss-of-function C. elegans mutants or by RNAi. Changes in cdr-1 transcription were then determined by measuring GFP expression or qRT-PCR. The expression of cdr-1, in both the absence and presence of Cd (100 μM), was suppressed when genes in the IIS pathway; daf-2, age-1, daf-18, pdk-1, akt-1, akt-2, sgk-1 and daf-16; were knocked down. Statistically significant interactions between Cd and IIS pathway genes were observed. Furthermore, knocked down of IIS pathway-related genes; skn-1, iso-1, pfa-4, pop-1, lin-14, tor-2, ras-1, vha-1; also inhibited cdr-1 expression in response to Cd. These results suggest that the IIS pathway mediates Cd-inducible transcription. In addition, they support a model where Cd exposure could induce elevated blood glucose levels by directly affecting the IIS pathway. The mechanism by which Cd activates the IIS pathway is currently being investigated.

**1158 Investigating a Protective Role of the Antioxidant N-Acetylcysteine against Cadmium-Induced Damage to Bone Extracellular Matrix.**


Environmental pollution of cadmium, a heavy metal commonly utilized in electronics, is a global human health concern. Bone is a known target site for cadmium. One mechanism to protect against cadmium-induced osteotoxicity and bone loss is by reducing oxidative stress. Our lab previously demonstrated that N-acetylcysteine (NAC) rescues osteoblasts from cadmium-induced apoptosis. Others have shown that reactive oxygen species generated through oxidative stress decreases bone mineral density, in part, through targeting the collagen component of the extracellular matrix (ECM). We hypothesize that the antioxidant NAC will protect against cadmium-induced damage to ECM produced by osteoblasts. Saso-2 cells were induced to mineralize and treated with 1mM NAC or 5μM CdCl2 only or in combination for 6 or more days. Upon termination cell viability, phosphate ECM deposition, and collagen secretion and distribution in the ECM were evaluated using MTT, von Kossa, SiriusTM, and immunofluorescence, respectively. Treatment with CdCl2 for 6 days resulted in increased phosphate deposition, which was restored to that of control in the presence of NAC. We also examined collagen, the main organic component of the ECM. When osteoblasts were induced to form an ECM, treatment with CdCl2 led to significantly less collagen secretion which was reflected in less collagen deposition into the ECM; these effects were reversed in the presence of NAC. Further, NAC and NAC combined with CdCl2 significantly increased cell viability compared to control, leading us to also investigate cell cycle progression. These studies provide further evidence that antioxidants promote bone health by reducing the toxic effects of cadmium. Research funded by NIH-INBRE P20 RR01654and P20 GM103408 and NIH R15ES05866 grants.

**1159 Cadmium Causes Injury to Pancreatic Islets That Is Associated with Caspase-3 Labeling.**

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Diabetes is a growing worldwide epidemic. There is increasing interest in how environmental contaminants can contribute to the onset of type II diabetes. Impaired insulin release is a hallmark of type I diabetes and is key in the progression of type II diabetes. Multiple epidemiological and experimental studies show that exposure to the metal cadmium (Cd), is associated with diabetes and reduced serum insulin. To examine the cytotoxic effects of Cd within pancreatic islets, male Sprague Dawley rats were injected subcutaneously with either saline (control) or Cd (0.6 mg Cd/kg/day, 5 days per week). After 6, 9 and 12 weeks of Cd treatment, pancreatic tissue samples were removed then fixed in formalin. Pancreata were sectioned and H&E stained to identify islets then examined for changes in islet histology. A trained veterinary pathologist scored each sample for cytotoxic signs and signs of necrosis and apoptosis. All pancreata from Cd treated animals had elevated scores for signs of vacuolization, apoptosis and necrosis. However, these changes in cell viability did not appear to change with longer Cd exposure times. In another study using the same pancreas samples, tissue was labeled for the apoptosis indicator, active caspase 3. In this study, pancreatic samples were counter stained with hematoxylin so that immuno-labeled islets could be identified. This study resulted in similar findings. Islets from Cd-treated animals had greater caspase-3 labeling and as before, the intensity of labeling appeared to be time independent. These preliminary results show that Cd acts to injury pancreatic islets which may result in diminished insulin release.

**1160 Regulation of Glutathione Synthesis in Cadmium-Treated Cultured Choroid Plexus.**

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In primary cultures of rat choroid plexus we have shown that exposure to cadmium (Cd) induces oxidative stress, stimulates apical choline uptake, and alters glutathione (GSH) synthesis. Our aim is to characterize the regulation of GSH availability and synthesis in response to low dose exposure to Cd and assess the significance of GSH in cellular adaptation to Cd. We treated choroid plexus primary cultures with 0 or 500 nM CdCl2 in serum free medium (SFM) for 12 h and collected samples at 3, 6, 9, and 12 h. Induction of heme oxygenase-1 (HO-1), heat shock protein 70 (HSP70), and metallothionein-1 (MT-1) in Cd-treated cells was compared to time-matched controls by immunoblot and qRT-PCR analyses. HO-1 protein was induced at 3 and gradually increased through 12 h. HO-1 gene expression was maximally induced by 70-fold at 6 h and was sustained through 12 h. HSP70 protein was maximally induced by 5-fold at 12 h. MT-1 gene expression was induced by 50-fold at 12 h. Cd induced the catalytic and modifier subunits of glutamate-cysteine ligase (GCL), the rate-limiting enzyme in GSH synthesis. GCLC protein levels peaked at 6 h while GCLM peaked at 9 h. Gene expression of GCLC and GCLM increased by 5-fold and 4-fold at 12 h. To elucidate the effects of Cd on GSH concentration, cells were pretreated (12 h) without or with buthionine sulfoximine (BSO, 100 μM) then treated with 0 or 250 nM CdCl2 ± BSO in SFM for 12 h. Intracellular GSH and oxidized glutathione (GSSG) concentrations were assayed and compared to control. Cd increased GSH by 2-fold and increased GSSG by 30-fold. In Cd-treated cells, BSO reduced GSH concentrations by 92% but increased GSSG concentrations by 15-fold above controls. Inhibiting GSH synthesis with BSO augmented induction of HO-1, HSP70, and MT-1 and enhanced stimulation of apical choline uptake in Cd-treated cells. These data indicate that Cd stimulates glutathione synthesis at the transcriptional level; and the inhibition of GSH synthesis accentuates cellular stress.

**1161 Stable Expression of FRET-Based Cd(2+)-Biosensor for Advanced Research of Cytotoxic Cd(2+).**

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Cadmium ion (Cd2+) causes lots of human tissue damages. Several ion channels and transporter proteins as the molecular gateway of different ions have been previously proposed to be the cytotoxic entry mechanism(s) of Cd2+. However, the solid conclusions have not been made. Here we tried to explore this issue through establishing human cell line stably expressing the fluorescent resonance energy transfer (FRET)-based Cd2+ biosensor for monitoring content of intracellular Cd2+. This newly constructed Cd2+ sensor Met-cad 1.57 contains part of Cd2+-binding protein CadR originated from bacteria Pseudomonas putida as the Cd2+ sensing key. Through G418 selection, the HEK293 line stably expressing Met-cad 1.57 HEK-MCD157 cells have been archived. The spectral patterns and sensing ranges of Met-cad 1.57 for intracellular Cd2+ sensing were characterized in vivo after 96-well plate. Under both a fluorescence spectroscopy and a FRET microscopic ratio imaging, the contents of Cd2+ were than monitored within living HEK-MCD157 cells. The dynamic range of Met-cad 1.57 from 0 to 100 μM of Cd2+ is 2–4.4 (2.2
fold) with a dissociation constant $K_d$ around 271 nM. The role of voltage-gated Ca2+ channels as the candidate of Cd2+ entry gateway was further explored with the HEK-MCD157 cells. In summary, a human embryonic kidney cell line HEK-MCD157 stably expressed FRET-based Cd2+ biosensor Met-cad 1.57 was constructed for both reliable and convenient investigations on content measurement of Cd2+ within live-cell including the identification of Cd2+ entry pathway and subcellular sequestration.

**1162 Blood Cadmium Level of Residents Living near the Abandoned Metal Mines in Korea.**

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Introduction: Preliminary biological exposure monitoring study selected 38 among 906 metal mines abandoned in Korea as the relative high risk areas for national surveillance subjects. They were surveyed from 2008 to 2011 under the environmental health policy by Ministry of Environment. The purpose of this study is to assess the blood cadmium levels from people living around a total of 38 abandoned metal mines and to determine if those areas have influenced exposure to cadmium at residents

Methods: Blood cadmium levels were quantified using graphite furnace atomic absorption spectrometry (AAAS) through appropriate blood collection, storage and pretreatment procedure. People living less than 3 km from the abandoned mine area were classified as the exposure group ($n=4,687$). Control group ($n=2,643$) was selected from people living in non-mining areas.

Results: Blood cadmium level from exposure group was found to be 1.59 ug/L as geometric mean (GM) (95% CI: 1.56 - 1.62 ug/L) and 1.91 ug/L as arithmetic mean (AM) (95% CI: 1.88 - 1.94 ug/L). Those levels were significantly higher than those from control group (GM=1.28 ug/L, 95% CI: 1.25 - 1.31 ug/L, AM=1.52 ug/L, 95% CI: 1.49 - 1.56 ug/L) ($p<0.0001$). In addition, 79.5% (n=3,728) of exposure group and 69.6% (n=1,840) of control group exceeded the reference value (1.0 ug/L) derived by German human biomonitoring commission. Multiple regression model found that exposure group was significantly higher than that of control group after age, gender and smoking status were all adjusted (adjusted $R^2=16.5\%$, $p<0.0001$)

Conclusions: We found that blood cadmium level of people residing near abandoned metal mine area was significantly high, compared with those of control group. Our findings suggested that the abandoned mine areas may have elevated blood cadmium level of people in the vicinity of those areas.

**1163 One-Week and Four-Week Inhalation Toxicity Studies of Nickel Sulfate and Nickel Subsulfide in Rats.**

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Inhalation toxicity studies of nickel sulfate (NiSO4) and Ni subsulfide (NiS2) were conducted in F344 rats to evaluate cellular pathway responses in lung distal airways among the candidate of Cd2+ entry gateway was further explored with the HEK-MCD157 cells. In summary, a human embryonic kidney cell line HEK-MCD157 stably expressed FRET-based Cd2+ biosensor Met-cad 1.57 was constructed for both reliable and convenient investigations on content measurement of Cd2+ within live-cell including the identification of Cd2+ entry pathway and subcellular sequestration.

Conclusions: We found that blood cadmium level of people residing near abandoned metal mine area was significantly high, compared with those of control group. Our findings suggested that the abandoned mine areas may have elevated blood cadmium level of people in the vicinity of those areas.

**1164 Gene Expression Profiles in Peripheral Blood Mononuclear Cells of Chinese Nickel Refinery Workers with High Exposures to Nickel and Control Subjects.**

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Background: Occupational exposure to nickel (Ni) is associated with an increased risk of lung and nasal cancers. Ni compounds exhibit weak mutagenic activity, after the cell’s epigenetic homeostasis, and activate signaling pathways. However, changes in gene expression associated with Ni exposure have been only investigated in vitro. This study was conducted in a Chinese population to determine whether occupational exposure to Ni was associated with differential gene expression profiles in the peripheral blood mononuclear cells (PBMCs) of Ni-refinery workers when compared to referents.

Methods: Eight Ni-refinery workers and ten referents were selected. PBMC RNA was extracted and gene expression profiling was performed using Affymetrix exon arrays. Differentially expressed genes between both groups were identified in a global analysis and findings were validated using real-time PCR.

Results: A total of 2167 differentially expressed genes displayed a greater than 1.25 fold-change in Ni-refinery workers. DNA repair and epithigenic genes were significantly overrepresented ($p<0.0002$). Of 31 DNA repair genes, 29 were repressed in the high exposure group and two were overexpressed while these genes were not repressed in the control group. Of the 16 epithigenic genes 12 were repressed in the high exposure group and 4 were overexpressed.

Conclusions: The results of this study indicate that occupational exposure to Ni is associated with alterations in gene expression profiles in PBMCs of subjects.

Impact: Gene expression may be useful in identifying patterns of deregulation that precede clinical identification of Ni-induced cancers.

**1165 Differential Severity of Chromium-Induced Hepatotoxicity in a Day.**

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The diurnal variation of cadmium-induced mortality is our topic (J Toxicol Sci, 37, 191, 2012). We presented further evidences for the Cd-induced diurnal variation and consider the mechanism of these in SOT 2012 meeting. In this study, we report the diurnal susceptibility of chromium-induced toxicity in mice. Male C57BL/6j mice adapted for 14 days with assigned to 6 groups of 5 animals were administrated intraperitoneally (i.p.) with hexavalent chromium (Cr(VI), 35 mg/kg) at different hours in the day (10:00, 14:00, 18:00, 22:00, 2:00 and 6:00 h), describing as zeitgeber time (ZT); ZT2, ZT6, ZT10, ZT14, ZT18 and ZT22. In case of dark period (ZT14, ZT18 and ZT22), administrations were performed under red light. The mortality was determined during 14 consecutive days. Mice were most sensitive to Cr administered at ZT10, while least sensitive at ZT22. In contrast, the susceptibility to Cd-induced mortality was higher at ZT6 but lower at ZT18. Therefore, the daily fluctuation of metal sensitivity differed depending on metal compounds, probably depending on the valence of metal ion. The GPT value, an index of hepatotoxicity, also showed higher at ZT10 injection of Cr. The hepatic levels of Cr determined by ICP-MS were not significantly different between both time (ZT10 and ZT22), thus Cr accumulation in the liver was not a determinator for this diurnal variation. Our results clearly indicate the susceptibility of metal-induced toxicity is markedly different in a day. We believe this viewpoint should be introduced into toxicology as “chronotoxicology” becoming indispensable in the future toxicology.

**1166 Speciation of Chromium Released from Metal-On-Metal Hip Implants.**

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Increased blood chromium (Cr) concentrations have been measured in many patients with hip prosthetic devices containing cobalt-chromium-molybdenum alloy. To date, no studies have attempted to analytically distinguish between non-toxic
trivalent Cr, Cr(VI), and potentially harmful hexavalent Cr, Cr(VI), in these patients. However, it is known that Cr(VI) preferentially accumulates in red blood cells (RBCs), while Cr(III) remains largely confined to the serum compartment of the blood. Hence, the relative distribution of Cr into RBCs and serum is often used as an indirect measure of Cr valence. We reviewed 231 blood samples from six different studies involving patients with Cr-containing hip implants. Samples were generally collected pre-implant and then at three or six month intervals post-implant for up to two years. The ratio of serum to RBC Cr concentrations was highly consistent. Post-implant serum Cr concentrations (median values of 0.17-2.39 μg/L) typically increased by several fold over pre-implant concentrations (median values of approximately 0.2 μg/L), but the RBC concentrations did not increase over time. Specifically, the RBC Cr concentrations did not change pre-vs. post-implant (median values of 0.26 to 2.5 μg/L). These data indicate that the Cr released from the implants is in the form of Cr(III).

DNA double strand breaks (DSB) are one of the most deleterious lesions that are induced by particulate chrome. If left un repaired or misrepaired, DSB can cause mutations or chromosomal aberrations leading to genomic instability. The aim of this study is to determine the genotoxicity of particulate chrome and investigate cellular responses of repair proteins after chronic exposure. We found chrome exposure induced concentration-dependent increases in DSB and with longer exposure time, the amount of breaks remained constant. To address whether this reflected a cycle of breakage and repair or deficient repair, we developed a human lung cell line that stably express GFP-53BP1 and generated time-lapse videos of chromosome treated cells. 53BP1 forms discrete nuclear foci at sites of DSB. By monitoring foci kinetics of these cells treated with chrome, we observed residual foci after 24 h, indicating some breaks persist presumably due to deficient repair. Therefore, we investigated the repair proteins involved in homologous recombinaton (HR) and non-homologous end joining (NHEJ) repair. We found that the nuclear fraction of phosphorylated ATM and Rad51 increased in 24 hours indicating HR is active. However, it largely decreased over time. We also observed concentra tion- and time-dependent decrease in Rad51 foci formation, which confirms that HR was inhibited. In contrast, protein expression of Ku80 and DNA-PKcs increased during chronic exposure to chrome. These results suggest that there is a signaling switch between HR and NHEJ during chronic exposure to chrome. Future work will focus on identifying key proteins that regulate the transitions between two repair pathways. This work was supported by NIHES grant ES016893 (J.P.W.) and the Maine Center for Toxicology and Environmental.

Cobalt (Co) ions are known to have a strong affinity for sulfhydryl proteins and amino acids in the blood and tissues, and sufficiently high concentrations of free ionic Co(II) can lead to adverse health effects in humans and animals. We describe a method using size exclusion liquid chromatography (SEC) to resolve protein-bound Co fractions from cyanocobalamin and free Co(II) after direct injection of serum samples stabilized with 0.1M acetic acid. Highly sensitive detection using ICP-MS was coupled with SEC to provide a method detection limit of 0.12 ng/L in human serum for one higher molecular weight protein-bound Co peak (likely Co(II) bound to human serum albumin (66 kDa) and other large serum proteins), as well as for cyanocobalamin (1.3 kDa) and for free Co(II). Validation studies show that this novel method demonstrates good accuracy with >98% mean recovery and 10.4% relative percent deviation. Good precision was demonstrated for matrix spikes at 0.5 μg Co/L with 92.7% mean recovery and 4% relative percent deviation. Other metal cations known to compete with Co(II) for protein binding sites (Fe(II), Zn(II), Mn(II), Cd(II), Cu(II), Ni(II), Pb(II)) did not significantly alter Co(II) quantitation in the stabilized acetate solution. The sum of protein-bound and free ionic Co species provided >90% mass balance when compared to total Co measurements analyzed by acid digestion and ICP-MS in the same human serum samples. We conclude that this method complies with the method performance criteria outlined by the U.S. Depart. of Health and Human Services guidance for industry bioanalytical method validation. These measurements focusing on identifying protein-bound and free Co(II) concentrations are likely to be more informative with respect to understanding interactions associated with adverse health effects in people with workplace Co exposures, dietary Co supplement users, and patients with elevated Co blood levels due to cobalt-chromium alloy prosthetic devices.

DNA double strand breaks (DSB) are one of the most deleterious lesions that are induced by particulate chrome. If left un repaired or misrepaired, DSB can cause mutations or chromosomal aberrations leading to genomic instability. The aim of this study is to determine the genotoxicity of particulate chrome and investigate cellular responses of repair proteins after chronic exposure. We found chrome exposure induced concentration-dependent increases in DSB and with longer exposure time, the amount of breaks remained constant. To address whether this reflected a cycle of breakage and repair or deficient repair, we developed a human lung cell line that stably express GFP-53BP1 and generated time-lapse videos of chromosome treated cells. 53BP1 forms discrete nuclear foci at sites of DSB. By monitoring foci kinetics of these cells treated with chrome, we observed residual foci after 24 h, indicating some breaks persist presumably due to deficient repair. Therefore, we investigated the repair proteins involved in homologous recombinaton (HR) and non-homologous end joining (NHEJ) repair. We found that the nuclear fraction of phosphorylated ATM and Rad51 increased in 24 hours indicating HR is active. However, it largely decreased over time. We also observed concentra tion- and time-dependent decrease in Rad51 foci formation, which confirms that HR was inhibited. In contrast, protein expression of Ku80 and DNA-PKcs increased during chronic exposure to chrome. These results suggest that there is a signaling switch between HR and NHEJ during chronic exposure to chrome. Future work will focus on identifying key proteins that regulate the transitions between two repair pathways. This work was supported by NIHES grant ES016893 (J.P.W.) and the Maine Center for Toxicology and Environmental.

One leading hypothesis for the carcinogenicity of hexavalent chromium (Cr(VI)) is that Cr(VI) causes a DNA double strand break (DSB) repair deficiency which leads to chromosome instability and neoplastic transformation. However, no studies have been done to determine if Cr(VI) can cause DSB deficiency in human lung cells. To begin to test this possibility, we exposed human lung cells to lead chrome for three sequential 24 h periods, each separated by a month. After each treatment, cells were seeded at colony forming density, cloned, expanded and retreted. Each generation of clones was tested for chromosome sensitivity, chromosome complement, DNA repair capacity and ability to grow in soft agar. We found that after the first treatment, lead chrome-treated cells exhibited a normal chromosome complement though a few clones showed a repair deficient phenotype. After the second exposure, more than half of the clones acquired an abnormal karyotype including numerical and structural alterations and many showed deficient DNA DSB repair. The third treatment resulted in more abnormal clones, previ ously abnormal clones acquiring additional abnormalities, and most clones were re pair deficient. Further investigation revealed that some clones were unable to form Rad51 foci in response to radiation, suggesting a defect in the homologous recom bination repair pathway. Finally, clones from the third generation were able to form colonies in soft agar suggesting that cell lines have been immortalized. This work was supported by NIHES grant ES016893 (J.P.W) and the Maine Center for Toxicology and Environmental Health.
Hexavalent chromium (Cr(VI)) is present in the marine environment and is a known human carcinogen and skin irritant. Cr(VI) is the form of chromium that is well absorbed through the cell membrane. It is also the most prevalent form in seawater. We compared the cytotoxicity of Cr(VI) in fin whale and human skin cells. Our data show that particulate chromium is cytotoxic to human and fin whale cells in a concentration-dependent manner. Specifically, data show that Cr(VI) is less cytotoxic to fin whale skin cells than human skin cells. For fin whale cells, we found that concentrations of 0.1, 1.0, 5, 10 and 20 μg/cm² lead chromate induced 91, 72, 55, 40 and 38 percent relative survival, respectively. In human cells we found that concentrations of 0.1, 0.5, 1.0, 5.0, and 10 μg/cm² lead chromate induced 64, 28, 10, and 0 percent relative survival, respectively. These data indicate that fin whale skin cells were 7.2 times more resistant to Cr(VI) than human skin cells. When corrected for intracellular ion uptake fin whale skin cells remained more resistant. We considered whether this effect was due solely to the chromium ion. Our data show that soluble chromium was also cytotoxic to human and fin whale cells in a concentration-dependent manner. They also indicate that fin whale skin cells were more resistant to soluble Cr(VI) than human skin cells. However, these soluble Cr(VI) differences were accounted for by intracellular ion uptake, which indicates that the difference cytotoxic effects are due to something other than the chromium ion by itself. Further investigations will look into the genotoxic effects of hexavalent chromium on these two species. This research is supported by the Environmental Protection Agency’s Undergraduate Research Opportunities Fellowship (CFW), NIEHS grant ES016893 (JPW) and the Maine Center for Toxicology and Environmental Health (JPW).

1174 Instability of Metallothionein Inhibits Mitochondrial Activities Following Exposure to Silver Nitrate in Human Bronchial Epithelial Cells.

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Silver (Ag) has been reported to generate reactive oxygen species (ROS) and to bind sulphhydryl groups in metabolic enzymes and non-enzymatic proteins such as metallothionein (MT). In the present study, we assessed toxicity of AgNO3 in human bronchial epithelial cells (BEAS-2B) focusing on the intracellular Ag distribution and ROS production. The cells were exposed to 0 – 10 μmol/l AgNO3 for 0 – 24 h and cytotoxicity was assessed with a modified MTT method. The cell viability was decreased by AgNO3 in a dose-dependent manner (IC50 = 2.5 μmol/l). Concentration of Ag in culture media decreased with time and stabilized at 12 h after exposure. Concentrations of Ag and Ag-bound MT in the soluble fraction of cells were sharply increased up to 3 h and then decreased, indicating that cytosolic Ag relocated to the insoluble fraction of cells shortly after Ag exposure. mRNA levels of major human MT isoforms, MT-I and MT-II, parallel concentrations of Ag, Cu and Zn-MT. The ROS-derived fluorescence intensity appeared to be elevated at mitochondria after treatment of the cells with Ag. To evaluate effects of Ag on mitochondrial respiration and activities of electron transport chain complex, mitochondria were obtained from rat liver tissues. The electron transport activities of all mitochondrial complexes (I to IV) were inhibited by Ag. We observed that a concentration of cytosolic H2O2 was increased up to 11.6 pmol per 10⁶ cells. Ag as well as Cu and Zn were released from isolated MT by H2O2 at concentrations as low as 0.3 μmol/l. Induction of Ag-stimulated H2O2 synthesis and Ag was suppressed immediately by MT. 3 h after exposure. MT was decomposed by cytosolic H2O2. Ag released from MT to insoluble cellular fractions and inhibited electron chain transfer of mitochondrial complexes, which eventually led to cell damage.

1175 Development and Optimization of a Procedure for the Determination of Indium-Tin Oxide Particle Size and Concentration in Cellular Media.

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Indium-tin oxide (ITO) is a solid mixture often comprised of approximately 90% indium oxide (In2O3) and 10% tin oxide (SnO2) by weight. ITO is employed as a transparent conductive coating for flat panel display and semiconductor devices and is typically deposited after sintering as a thin film on the desired substrate through a variety of technologies. Fatal cases of interstitial pneumonia and alveolar proteinosis have been reported for workers exposed to ITO particles. In vitro studies of ITO are planned in order to better understand the toxicity of this compound. Comprehensive characterization of ITO test materials is required prior to toxicity testing. Characterization of ITO particle size under the conditions of laboratory testing is important because this physicochemical parameter can significantly impact bioavailability and cellular toxicity.
In these studies we developed, optimized, and applied a dynamic light scattering (DLS) sample preparation and measurement protocol for determining hydrodynamic particle size in a suite of sintered/non-sintered ITO samples. ITO samples were prepared in cellular growth media at doses ranging from 0.3 – 0.4 mg/mL. Sonication times of particle suspensions were evaluated from 15 – 90 minutes. Times ranging from 30 – 60 minutes yielded the most stable suspensions with respect to hydrodynamic particle size over a 24-hour period. Resulting suspensions from samples with 30 – 60 minute sonication times were stable for 10 – 24 hours after sonication, with respect to hydrodynamic particle size. Throughout the 24-hour DLS measurement period, suspensions were analyzed for indium content by inductively coupled plasma mass spectrometry (ICP-MS) to confirm ITO concentrations. The developed protocol enabled investigators to use ITO suspensions of known particle size and concentration in their in vitro cellular studies.

**1178 ICP-MS Determination of Tissue Iron Levels in the Rat.**

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Iron is an essential trace element associated with toxicities when present in excess, as for example in conditions such as hereditary hemochromatosis. We have developed and validated a reliable and rapid quantitative method for the exploration of endogenous levels of total iron in rat plasma, heart, liver, spleen, kidney, stomach, lung and bone marrow. The analysis by ICP-MS (Inductively Coupled Plasma Mass Spectrometry) was selected among all other techniques, because of its easy handling of simple or complex matrices requiring very little sample preparation. The method was evaluated for accuracy, precision, linearity, matrix effect and instability derived from adsorption on container walls. Precision and accuracy for all matrices ranged from >4.3% to <7.9% and >1.5% to <12.2% respectively. Matrix effect was negligible and the assays for the evaluation of instability showed no adsorption of iron on container walls, either in glass, polypropylene or polyethylene. Iron levels were determined over a 28-day period, on 145 rats aged 7 to 8 weeks at the beginning of the study. In general a low to moderate inter-animal variability of endogenous iron levels was observed. The levels of total iron in plasma, heart, liver, stomach and bone marrow were stable over time with concentrations <5000 ng/mL in plasma and <200 ng/g in the other tissues. In conclusion, a reliable and rapid ICP-MS method was used to determine endogenous levels of total iron in rat plasma and tissues. This method can readily be adapted for the study of endogenous iron levels in other laboratory animal species.

**1177 Genetic Variation of Iron Metabolism in Mice.**

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Iron deficiency is the most common disease in the world with an estimated 4.5 billion affected persons. Debilitating fatigue, altered immune function, decreased work capacity and anemia are among the deleterious consequences of this pervasive disorder. Similarly, liver iron overload is a significant health concern. Multiple genetic disorders of iron metabolism in man, rodents and other vertebrates suggest multiple loci can contribute to the susceptibility of iron deficiency and severity of iron overload. Previous studies have shown genetic variation in iron metabolism between inbred strains of mice. We hypothesize that this genetic variation underlies the differences seen in iron metabolism between inbred mice. Our aim is to map the quantitative trait loci responsible for the strain specific differences in iron metabolism. To do this we will use high resolution SNP analysis to “in silico” map the quantitative trait loci responsible for the divergent iron related phenotypes using the hybrid mouse diversity panel (HMDP) [1]. Here we show the phenotypic analysis of the variation in iron metabolism for these mice, and how the overall elemental profiles change for each strain.

References

**1176 Tungsten Exposure Increases Lung Metastases in an Orthotopic Murine Breast Cancer Model.**

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Tungsten is a strong, flexible metal that until recently had been thought to be an “inert” metal. These properties led to its incorporation into the manufacturing of medical devices. In a recent clinical trial, a tungsten-based shield was used in the treatment of breast cancer patients undergoing intraoperative radiotherapy. Following the procedure, the women were left with residual tungsten in their breasts. Elevated tungsten levels in the blood and urine indicate that tungsten is not remaining in the breast tissue. Tungsten was detected in the urine of patient even 8 months after mastectomy, indicating another reservoir has been created. Based on previous data, we hypothesize this reservoir to be the bone. Animal studies suggest that tungsten may contribute to carcinogenesis and can alter development and increase DNA damage in the immune system. In order to evaluate the effect of tungsten on breast cancer, female Balb/C mice were exposed to 15 ppm of sodium tungstate for 1 month followed by injection of 6000 cancer cells into the mammary fat pad. The size of primary tumor, the extent of lung metastases and immune parameters were evaluated. Tungsten did not alter the growth of the primary tumor. However, the number and average size of lung metastases was significantly greater in the tungsten-exposed animals. This model is not known to metastasize to the bone, but we found that tumor-bearing mice had 3-fold more tungsten in the bone than non-tumor bearing mice. Tungsten increased the peripheral blood leukocyte count in non-tumor bearing mice, but decreased the massive granulocytosis associated with tumor growth. These data suggest that tungsten increases the extent of breast cancer metastasis to the lung, which could have a significant impact on individuals who have cancer and are also exposed to tungsten. The levels of tungsten deposition within the bone and immune cell parameters were also altered, which could also impact metastasis in this model.

**1179 Development of a Transportable Neutron Activation Analysis System to Quantify Manganese in Bone In Vivo—Feasibility and Methodology.**


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Manganese (Mn) is a vital element in human body for growth and other functions. However, overdose to Mn compounds can lead to many adverse health effects. An important issue in assessing Mn exposure is to select a proper biomarker. Over 40% of Mn in human body is stored in bone and bone Mn represents long term chronic Mn exposure. Hence it is attractive to develop a technology to quantify the concentration of Mn in human bones in vivo. Neutron activation analysis (IVNAA) is a promising non-invasive technique which can determine the concentrations of various elements in the human body. In our study, we investigated the possibility and feasibility of developing a transportable IVNAA system for Mn quantification in human bone. Monte Carlo simulations using a portable deuterium-deuterium (DD) neutron generator as neutron source. Experiments were conducted with a deuterium-tritium (DT) generator available in our lab to validate the MC simulation results. The count rates calculated from MC simulations were in agreement with a deuterium-tritium (DT) generator available in our lab to validate the MC simulation results. The count rates calculated from MC simulations were in agreement with those obtained from the experiments for the DT generator system setup. Different types of moderator and reflector for DD setups were simulated and paraffin was selected as the best material for both of the moderator and reflector. Then the optimal thicknesses for the reflector and moderator were determined. Assuming normal concentration of Mn in bone is about 1 μg/g and neutron yield is 4x10^6/sec, the count rate is 0.324 counts/sec if we use a 4π NaI detector with 100% efficiency. The corresponding dose to the hand for 10 minutes irradiation time is about 43.704 mSv from MC simulation, which corresponds to about 48.64 μSv total body equivalent dose. In conclusion, it is feasible to develop a transportable NAA system to quantify Mn in bone in vivo with a minimum radiation dose exposure to the subject.
develop analytical means to measure each isoform of this diverse protein family. The twelve human MT isoforms share 70 to 90% amino acid sequence identity which hampers antibody-based methods of isoform-specific quantification at the protein level. Trypsin digestion yields an N-terminal MT peptide that has a unique mass for each MT isoform. Each of these peptides contains five Cys residues, is N-terminally acetylated, and once alkylated is relatively hydrophilic. The human kidney HK-2 epithelial cell line expressing stably transfected MT-3 was used as a model system. Cytosol was prepared from control cells and cells exposed to 9 μM cadmium for 3 days. The cytosol was then denatured with urea, reduced with DTT, and alkylated with either 14N- or 15N- iodoacetamide. For absolute quantification, -145 pmol of MT isoform-specific 15N-labeled N-terminal acetylated peptide was added to the samples. The light and heavy-labeled samples were combined and digested with trypsin. Strong cation exchange (SCX) and C18 reverse-phase liquid chromatography (rp-HPLC) successively enriched the N-terminal MT peptides followed by MALDI-TOF/TOF MS and MS/MS. The ratio of light (control) vs. heavy (induced) MT peak intensities were calculated for each MT isoform detectable in both the control and Cd-induced samples. Seven MT isoforms were detected and quantified in the HK-2-MT3 cells. MT-1F showed the largest induction (10-fold) from 9 μM Cd treatment. Metallothionein 3 was induced 2-fold. This study describes an iodoacetamide stable isotope-labeling mass spectrometry-based method for simultaneous relative and absolute quantification of human metallothionein isoforms.

1181 Identification of Metallothionein-3 Interacting Proteins in the Human Proximal Tubule Cells.

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Metallothionein 3 (MT-3) is a small, cysteine-rich protein that binds to essential metal ions required for normal cellular processes, as well as heavy metals that have the potential to exert toxic effects on cells. MT-3 is expressed by epithelial cells of the human kidney, including the cells of the proximal tubule. This laboratory has previously shown that the mortal cultures of human proximal tubule (HPT) cells express MT-3 and could form domes in the cell monolayer, a morphological feature indicative of vectorial active transport. However the immortalized proximal tubular cell line HK-2 lacked the expression of MT-3 and failed to form domes. Transfection of the HK-2 cells with the MT-3 gene restored dome formation in the cells suggesting that MT-3 may have a role in vectorial active transport. Recent in vivo and in vitro studies have reported that MT-3 is interacting with other proteins and these interactions are thought to be playing an important role in execution of some of its functions. Thus the goal of this study was to identify the binding partners of MT-3, which would allow us to understand the mechanism through which MT-3 is regulating vectorial active transport and cell differentiation in HPT cells. For this purpose cell extracts were prepared using the immortalized human proximal tubular (HK-2) cell line. Protein pull-downs were then followed by SDS-PAGE and mass spectrometry analysis. Data analysis revealed an interaction between MT-3 and myosin 9, gelsolin, beta-actin, and tropomyosin 3. Immunofluorescence studies further confirmed that MT-3 co-localizes with these proteins. These studies demonstrate that MT-3 is interacting with proteins that are involved in cytoskeleton organization of the cell and thereby regulating vectorial active transport and cell differentiation.

1182 The Unique N and C-Terminal Domains of Metallothionein-3 Influence the Growth and Differentiation of MCF-7 Breast Cancer Cells.

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Toxic insult from the heavy metal cadmium (Cd²⁺) is known to induce the expression metallothioneins (MT) which are heavy metal binding proteins. Previous work from our laboratory has shown that over-expression of MT-3 in breast cancers is associated with poor patient outcome. Furthermore, MT-3 has been shown to inhibit the growth of breast cancer and prostate cancer cell lines. Studies have shown that the MT-3 protein contains 7 additional amino acids that are present in any other member of the MT gene family, a 6 amino acid C-terminal sequence and a Thr in the N-terminal region. The unique N-terminal sequence is responsible for the growth inhibitory activity of MT-3 in the neuronal system. The goal of this study was to characterize the function of the N and C-terminal domains of MT-3 in the breast cancer cell line, MCF-7. For this purpose six different constructs of MTs were prepared which were as follows: wild type (WT) MT-3, MT-3 N-terminal deletion (MT-3 NT), MT-3 C-terminal deletion (MT-3 CT), WT MT-1E, and MT-1E mutated to contain the N-terminal of MT-3 or the C-terminal or both the N- and the C-terminal of MT-3. Each of these constructs was transfected into MCF-7 cells and the growth rates and the transepithelial resistance (TER) was measured. The data obtained suggests that the N-terminal region of MT-3 is involved in growth inhibitory activity whereas the C-terminal region is involved in vectorial active transport which is indicated by the formation of domes in cell culture and an increase in TER. In conclusion, this study further characterizes the unique properties of metallothionein MT-3 and the potential role that it may play in the differentiation of certain breast cancers.

1183 Fluoride, Aluminum, and Mixtures: Histological Changes in the Liver of Mouse, Mus norvegicus albinus.


Fluoride is one of the chemical contaminants of water causing several health problems in man and animals. Aluminum is one of the most abundant metals in the environment. Although aluminum has been historically considered as a non toxic metal, which is toxic in its trivalent (Al³⁺) form. Earlier studies exhibited fluorosis and aluminiumosis when fluoride and aluminium exposed individually. However, it is important to study two different toxicants in combination to know whether they can act as synergistic or antagonistic. In this study the toxic effects exerted by the accumulation of fluoride, aluminum and their mixtures were studied in the liver of mice. Six different doses were tested for a period of 30 days and 60 days for their toxic effect in the liver. Microscopic study of liver tissue revealed interesting pathological changes in all the fluoride treated mice. The damage to the tissue was dose and time dependent. The aluminum treated mice showed conspicuous histopathological changes in higher dose, which was more significant with the increase in period of exposure. Whereas, the histopathological changes, were not much conspicuous in lower dose groups, and also decreased with period of exposure. In the mixture of fluoride and aluminum treated mice, mild to severe changes were observed in the higher dose of fluoride with aluminum, whereas very mild and insignificant changes were observed in the mice exposed to lower dose of fluoride with aluminum.

1184 Both the Metal Moiety and the Organic Backbone of Ethylene Bisdithiocarbamate Fungicides Maneb and Mancozeb Contribute to Toxicity in Human Colon Cells.

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Ethylene bisdithiocarbamate (EBDC) pesticides are used primarily as broad range contact fungicides on a wide variety of crops. A subset of the metal containing EBDC pesticides include Maneb (MB), Mancozeb (MZ), and Zineb (ZB), which are complexed with the transition metals manganese, zinc, or in the case of MZ, both manganese and zinc. While these agents are reported to possess low human toxicity, previous testing in our laboratory has established the toxicity of these compounds to transformed colon cells, HT-29 and Caco2, and to normal colon cells, CCD-18Co. Significant decreases in viability were observed with MB and MZ in HT-29 and CCD-18Co (80-260μM), and Caco2 cells (40-180μM). ZB exposure however, produced no significant decrease in cell viability in all cell types up to 800μM. Therefore, the purpose of this study was to determine if the metal moieties of MB and MZ contributes to the generation of toxicity. To determine if the manganese and zinc can accumulate within HT-29 and Caco2 cells and initiate toxicity, inductively coupled plasma-optical emission spectroscopy was performed. MB exposure resulted in a dose dependent increase in intracellular manganese levels within HT-29 and Caco2 cells (20-200μM), MZ exposure also resulted in a significant accumulation of zinc within HT-29 and Caco2 cells (40-200μM). To investigate whether manganese and/or zinc was a critical factor in the cytotoxic process, HT-29, Caco2, and CCD-18Co cells were treated with manganese chloride (MnCl₂) and zinc chloride (ZnCl₂). MnCl₂ and ZnCl₂ exposure produced no significant loss of viability in all cell types up to 400μM. The lack of toxicity observed following treatment with ZB within the same concentration range as MB and MZ suggests that the manganese portion of these agents may play a key role in generating the effects seen upon exposure. However, the lack of differences in the cytotoxicity of MnCl₂ and ZnCl₂ treated cells indicates that the EBDC backbone may act in conjunction with the metal moiety to cause toxicity.
Inhalation of metals is associated with development and aggravation of lung and cardiovascular diseases, and inflammation is considered to be a central pathological mechanism. In this study we have investigated expression of inflammation-related genes in human bronchial epithelial cells (BEAS-2B) exposed to cadmium (Cd), zinc (Zn), arsenic (As), vanadium (V), manganese (Mn) and nickel (Ni), metals often found in polluted air. As measured by real-time PCR array, Cd, Zn and As induced cytokine expression patterns differing from V, Mn and Ni, with higher levels of CXCL8, CXCL5, IL-6 and CCL26. Furthermore, Cd, Zn and As, but not V, Mn and Ni, induced increased levels of 1-heme oxygenase indicative of oxidative stress. Thus, the mechanisms of Cd-, Zn- and As-induced cytokine responses were investigated further. In line with the gene-expression data, Cd, Zn and As also induced high levels of CXCL8 and IL-6 protein-release, and addition of antioxidants reduced the metal-induced cytokine responses. Moreover, the metals activated the mitogen activated protein kinases (MAPK) p38, JNK and ERK and the p65 subunit of NF-kB, measured as protein-phosphorylation by Western blotting after 2 and 4 hr. Use of MAPK-inhibitors and siRNAs against p65 revealed that these signaling pathways were involved in the cytokine responses. The p38 pathway seemed most important for both IL-6 and CXCL8 release induced by all three metals, whereas JNK seemed to be mainly involved in As-induced CXCL8 and ERK in As-induced IL-6 and CXCL8 releases. In conclusion, among the tested metals Cd, Zn, As were potent inducers of inflammatory responses and oxidative stress in epithelial lung cells. NF-kB and p38 signaling seemed central to metal-induced IL-6 and CXCL8 responses, whereas the relative involvement of JNK and ERK differed depending on the metal and cytokine.

Metallothioneins (MTs) are highly conserved, cysteine-rich metal-binding proteins that function in metal homeostasis and detoxification, as well as free-radical scavenging. To better understand the role of MT in cadmium-associated cellular responses, regulatory factors and pathways that control the expression of the C. elegans MT gene, mtl-1, were identified through genetic screens. A mutagenesis screen identified a mutant, JF99 that harbored a mutation affecting atf-7, a negative transcription factor involved in the JNK/p38 pathway. In both JF99 and an atf-7 deletion mutant, steady-state mtl-1 mRNA levels were identical but significantly greater than levels in wild type nematodes in the absence of cadmium. In addition to atf-7, mutations in pmk-1 caused a decrease in mtl-1 expression, suggesting that PMK-1 regulated ATF-7 activity. A candidate gene screen identified three insulin signaling pathway genes (PDK-1 and the AKT-1/AKT-2 complex) that functioned independently of this pathway. Further genetic analysis confirmed that these genes act upstream of PMK-1 and ATF-7 to regulate mtl-1 transcription. Based on our genetic pathway data and previous work, we propose that ATF-7 resides on the promoter region of mtl-1 to inhibit the constitutively active transcription factor ELT-2, which is important for intestinal cell specific transcription. In the presence of cadmium, upstream factors signal PDK-1 and the AKT-1/2 complex to phosphorylate PMK-1 causing it to translocate to the nucleus and phosphorylate ATF-7, allowing ELT-2 to initiate transcription. The activation of this pathway results in an increase in MT, which scavenges cadmium and free radicals. In mammalian cells, knockdown of PDK1 and ATF7 resulted in changes in MT-1 expression, suggesting that this pathway was not unique to C. elegans. The insulin signaling pathway affects the aging process, thus an association between cadmium exposure and the activation of portions of the insulin signaling pathway provides a mechanistic link between metal homeostasis, ROS and aging.

The transcription factor fos related antigen-1 (fosl1) is an AP-1 protein whose expression is often elevated in tumors and transformed cells. Previous studies show that fosl1 is transcriptionally regulated through the activity of the fos transcription factor and the MAPK pathway. In addition, fosl1 is induced after exposure to cadmium; however, the mechanism by which this induction occurs has not been elucidated. Metals, such as cadmium, are able to induce the expression of a wide variety of genes through the activation of metal-regulatory transcription factor 1 (MTF-1). Upon activation, MTF-1 translocates to the nucleus and binds to metal responsive elements (MREs) in the promoters of metal-inducible genes. To determine if MTF-1 participates in cadmium-inducible fosl1 regulation, a five kilobase region upstream of the transcription start site and the coding region of fosl1 were analyzed for MRE sequences. Seven MREs were identified and chromatin immunoprecipitation (ChiP) analysis performed to determine if MTF-1 had the ability to bind the fosl1 promoter after metal exposure. ChiP assays showed that MTF-1 localized to the fosl1 promoter and bound each MRE sequence to varying degrees. To assess the biological impact of this binding, MTF-1 null (dko7) cells and dko7 cells transformed with MTF-1 (dko7 + MTF-1) were exposed to cadmium and RNA was isolated at various time periods to evaluate steady state mRNA levels of fosl1 by real-time PCR (RT-PCR). These studies showed a biphasic increase in fosl1 mRNA in dko7 + MTF-1 cells up to 12 hours after exposure to 5 µM cadmium after which the mRNA levels declined to background levels at 24 h. In contrast, dko7 cells showed a steady increase in fosl1 mRNA regulation for the duration of the study. These data suggest that MTF-1 is recruited to the fosl1 promoter and may be a negative regulator of fosl1 mRNA expression.

Organophosphate (OP) pesticides represent a threat to soldiers, first responders and other civilians since they are highly toxic and widely accessible chemicals used in crop, industrial, and home applications. However, for many OP pesticides such as MP, there is incomplete toxicological data in humans. Most of the reports are epidemiological studies, adult and juvenile animal studies, or in vitro studies using transformed cell lines. Since undifferentiated human embryonic stem cells (hESC) maintain the ability to differentiate into any somatic cell in the body, they offer a unique vantage point to measure the effect of toxics on the very early human development and cell repair system. From our earlier metabolomics work examining the secretome of hESCs following exposure to MP, we hypothesized that ROS is generated during the toxic response to MP. In this follow-on work, we chose to visualize and quantitate: 1) the presence
and location of ROS, and 2) cell and nuclear morphology in MP-exposed WA09 hESC using the Cellomics Oxidative Stress I Assay on a Thermo Scientific Cellomics® ArrayScan® HCA System. While low levels of oxidative stress are necessary for normal cell metabolism, high levels are associated with cellular dysregulation and pathologies such as cancer. As expected, our imaging data indicated that the cells did not change structurally, nor did they undergo senescence or apoptosis in response to the MP exposures examined. While we were able to measure cellular oxidative stress (fluorescent conversion of dihydroethidium, DHE), the amount measured in the nuclei was much less than we had predicted. All other DHE was scattered and punctuate throughout the cell. These data, combined with other studies suggesting that hESC lose their robust ROS clearance protection mechanism as they become more differentiated, opens up a new avenue of investigation to define the point in embryonic differentiation when cells become more vulnerable to the ROS-induced toxic effects of MP.

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OCT4, SOX2, and NANOG are transcription factors that maintain pluripotency and self-renewal of embryonic stem cells (ESC) in culture: on the one hand, they activate expression of self-renewal genes, and on the other hand they prevent differentiation by silencing the expression of developmental genes. Unlike DNA methylation, polycomb group (PcG) proteins are chromatin modulators that repress the expression of developmentally important genes, such as those in the Hox family. PcG-mediated repression represents a plastic and dynamic way to keep genes poised to be reactivated during development. Promoters of developmental genes in ESC are often bivalently marked by trimethylation on lysine-4 (H3K4me3) and lysine-27 (H3K27me3) of histone H3, with PcG-mediated repression allowing these genes to be ready for reactivation upon differentiation. The aryl hydrocarbon receptor (AHR) has long been believed to mediate xenobiotic toxicity; increasing evidence suggests that it also plays a role during embryogenesis. To study AHR functions during early development, we have analyzed the regulation of AHR expression in C57Bl/6 mouse ESC and their in vitro differentiated progeny. In ESC, Ahr expression is silenced through bivalent marks in its promoter. Specifically, H3K27 is trimethylated in both distal and proximal promoter regions (dpr and ppr) and H3K4 is trimethylated in the ppr. We also detect the specific binding of pluripotency factors and PcG proteins in the dpr. RNA polymerase II appears to be paused in the ppr, producing short aborted Ahr transcripts. Upon differentiation, the Ahr promoter resolves into a H3K4me3 monovalent state and recruits Sp factors, leading to transcription activation. These results define an ESC-specific silencer domain located 2kb upstream of the Ahr gene transcription start site, responsible for the low, if any, Ahr expression in ES cells. Directed silencing, resulting from cooperation between pluripotency and PcG-repressive factors suggests that the Ahr may play a significant role in maintenance of pluripotency. Supported by NIH Grant 5R01ES06273

YSU, E. I. Tokar, B. J. Person, O. Orihuela and M. P. Waalkes, NTP, NIEHS, NIH, Research Triangle Park, NC.

Cancer stem cells (CSCs) are likely key to carcinogenesis. In prior work we found that arsenic-transformed malignant epithelial cells (MECs) recruit nearby, but noncontiguous normal stem cells (NSCs) into a CSC-like phenotype. Here, we tested if this recruiting effect on NSCs occurs with MECs transformed by other carcinogens. Normal human prostate epithelial SGs (WPE-stem) were co-cultured via transwell dishes that separate cells but not secreted factors with isogenic MECs derived from normal human prostate epithelial RWPE-1 cells by cadmium or N-methyl-N-nitrosourea (MINO) exposure (called Cd-MECs or MNU-MECs, respectively). SGs co-cultured with normal RWPE-1 cells served as control. After 2 weeks of Cd-MEC co-culture, SGs showed high secretion of metalloproteinase-9 (MMP-9, 3.6-fold of control) and MMP-2 (2.2-fold), increased colony formation (1.8-fold) and invasion (2.5-fold), formation of aggressive ductal structures in MECs, and loss of tumor suppressor gene, PTEN, expression (62% of control), all indicative of cancer phenotype. MNU-MEC co-culture had no such effects on SGs. Epithelial-to-mesenchymal transition occurred in Cd-MEC co-cultured SGs, as shown by cell morphology and VIMENTIN over-expression, but not in MNU-MEC co-cultured SGs. During Cd-MEC co-culture, loss and regain of SC-related gene expression (ABCG-2, OCT-4 and WNT7B) occurred, likely as a key to MGMEC recruitment of NSCs into CSCs. Overall, the data indicate SGs gain a cancer phenotype with Cd-MEC co-culture but not with MNU-MECs. Analysis showed Cd-MECs secreted high levels of the tumor microenvironmental related factor, TGFβ-1. Direct TGFβ-1 treatment of NSCs duplicated most responses from Cd-MEC co-culture. Since recruitment of NSCs into CSCs can also occur with arsenic-transformed MECs, we conclude that the effects of nearby MECs on NSCs are not carcinogen specific but apply to some, but not all, carcinogens. This phenomenon may be important in chemically-induced tumor extension and dissemination.

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Inorganic arsenic (As) is a human carcinogen probably targeting the prostate and can malignantly transform the human prostate epithelial cell line, RWPE-1 (to CAE-PE cells) and a derived stem cell (SC) line, WPE-stem (to As-ESC cells) by unclear mechanisms. MicroRNAs (miRNA) are non-coding but exert negative control on expression by degradation or translational repression of target mRNAs. Aberrant miRNA expression is likely a key factor in carcinogenesis. By miRNA array of 84 miRNAs in CAE-PE and As-ESC cells, for 29 and 13 miRNAs respectively, expression changed by >1.5 fold of control, and 7 were common in both cell lines. Increased miR-143 (5 fold), miR-205 (4 fold), miR-181d (3 fold), and miR-181c (2 fold) in CAE-PE cells correlated with increased RAS oncogenes, RAS, RAS2A, RAS2B, miR-205 and KRAS protein. Reduced miR-134 (5 fold), miR-134c-5p (3 fold), and miR-205 (2 fold) in As-ESC cells correlated with increased target RAS mRNA and, KRAS, NRAS, and RRAS proteins. The RAS/ERK signaling pathway helps control cell survival, differentiation, and proliferation, and if
1194 Aryl Hydrocarbon Receptor Contributes to Mouse Embryonic Stem Cell-Derived Cardiomyocyte Differentiation.
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2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) is a ubiquitous toxicant persistent in the environment. It is a prototypic ligand for the aryl hydrocarbon receptor (AHR), a ligand-activated transcription factor whose activation upon ligand binding regulates the expression of many xenobiotic detoxification genes. Besides being a critical mediator of gene-environment interactions, AHR has functions beyond those of a xenobiotic interacting protein. Studies in Ahr⁻/⁻ mice have shown that AHR has physiological roles in hematopoiesis and heart development. To characterize these roles and the consequences of TCDD exposure in cardiomyocyte specification, we studied the effects of TCDD exposure during mouse embryonic stem cells (mESCs) differentiation into cardiomyocytes. mESCs were allowed to spontaneously differentiate via embryo-like aggregates (embryoid bodies, EB) into contracting cardomyocytes. Immunocytochemistry on contracting EBs indicates that AHR colocalizes with the cardiac markers cardiac troponins Ⅰ and Ⅳ and NKX2.5. ShRNA knockdown of AHR during differentiation or TCDD treatment significantly decreased the percentage of contracting EBs. A time series of TCDD treatment revealed that the time between day 0 and day 3 of mESC differentiation was the most critical window for TCDD toxicity. An ESC clone containing a stably integrated plasmid bearing a Cyp1a1 promoter-driven puromycin-IRES-EGFP was allowed to differentiate in the presence of TCDD followed by puromycin selection. Real-time PCR analyses of the puromycin resistant selected cells showed that many mesoderm marker genes were up-regulated while both endoderm and ectoderm markers were down-regulated, indicating that early AHR activation might be committed to mesodermal lineages. Our data suggest that AHR is involved in the regulation of cardiomyocyte differentiation during embryonic development and that TCDD exposure during early differentiation is toxic to cardiomyocyte development. Supported by NIH Grant 5R01ES06273.

1195 Exposure to Concentrated Air Particles Depletes Bone Marrow Hematopoietic Stem Cells.
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Results from several epidemiological and laboratory studies suggest that exposure to airborne particulate matter (PM) is associated with an increase in cardiovascular disease (CVD) risk. While acute PM exposures have been linked to arrhythmias, thrombosis, inflammation, endothelial dysfunction, myocardial infarction and stroke, long term exposures increase the risk of mortality due to heart failure, cardiac arrest, and ischemic heart disease arrhythmias. Despite these associations however, relatively little is known about the molecular mechanisms linking exposures to CVD risk. In previous studies we have shown that circulating endothelial progenitor cells (EPCs) were depleted in humans exposed to natural variations in fine particulate matter (PM2.5) and in mice exposed to concentrated air particles (CAPS). However the effects of exposure on other progenitor cell types remains unknown. Hence, to examine how PM affects other stem cell populations we exposed mice to CAPS generated from downtown Louisville, Kentucky using a versatile aerosol concentration emission system (VACES) for 9 days and examined levels of bone marrow-derived hematopoietic stem cells (HSCs) and mesenchymal stem cells (MSCs) using colony forming assays. In CAPS-exposed mice we observed an approximate 35% decrease in CFU-GM (colony forming unit-granulocyte, macrophage) colonies compared to filtered air-exposed, control mice. However, the absolute number of colonies and the relative decrease versus air controls varied directly with CAPS levels. Flow cytometry-based phenotypic analysis of these colonies suggested a uniform decrease of multiple hematopoietic lineages rather than a decrease in a specific subset. In contrast no change was observed in BFU-E (burst forming unit-erythroid) colonies between the filtered air and CAPS-treated mice, or in the number of bone marrow-derived MSCs. These results suggest that HSC depletion may contribute to the cardiovascular complications resulting from chronic PM exposure by affecting immune responses to injury and dysfunction.

1196 Cadmium Concurrently Affects Cell Cycle, Total Histone Protein Production (THP) and H3 Histone Modification Pathways in Mouse Embryonic Stem Cells.
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The fetal basis of adult disease (FEBAD) theorizes that embryonic challenges initiate pathological in adult life through epigenetic modification of gene expression. We previously reported (2012) that cadmium (Cd) exerts differential toxicity in mouse embryonic stem cells (mESCs) by targeting THP production and H3 (a core histone protein) early in stem cell development, while H3K27 mono-methylation (H3K27me1) associated with transcriptional activation is affected in later stages of differentiation. Thus, low dose acute Cd exposure selectively disrupts chromatin structure, an effect not seen in differentially matured cells. In this study we tested the effects of Cd on cell cycle progression (flow cytometry), chromatin structure and epigenetic pathways (THP and H3K27me1 analysis, respectively) in undifferentiated mESC after 1-h and 24-h exposures plus recovery. The data suggest that mESC do not recover from Cd insult at 0.03 mM (IC50 for MTT assay), whereas cells recovered from 1-h exposure at 0.01 mM (IC25 for MTT assay). This confirms our previous results that maximum cytotoxicity is seen during the first few hours of exposure at low concentrations. Additionally, THP production is suppressed to a greater extent than cell proliferation at 0.01 mM (24-h). Interestingly, 1-h exposure plus recovery resulted in a significant increase in THP production. Furthermore, 24-h exposure (with recovery) suppressed both cell proliferation and THP production, indicating that Cd targets THP production at longer exposure times and disrupts repair mechanisms. Flow cytometry data demonstrates that 0.01 mM Cd, for 24- and 48-h, increases DNA synthesis and percent of cells in G1 phase as well as percent of cells in mitosis; differentiation is affected at 72-h only. Thus presently we report that low dose acute Cd toxicity is cumulative and disrupts DNA repair, while concurrently affecting cell cycle progression, repair mechanisms, chromatin structure and transcriptional state in mESCs.

1197 Cadmium Induces Up-Regulation of Genes Associated with Early Embryonic Development in Mouse Embryonic Stem Cells.
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Cadmium (Cd) alters gene expression and increases cell proliferation, but the exact mechanism of its nongenotoxic effect has yet to be elucidated. Previously we reported that exposure to low concentrations of Cd (80 nM to 5 μM) causes a toxicologically relevant increase in cell proliferation, as determined by [H]-thymidine incorporation, in undifferentiated pluripotent mouse embryonic stem cells (mESCs). Since epigenetic mechanisms are important during early embryonic development, we evaluated the potential of Cd to disrupt normal embryonic processes in these cells. Using trypan blue exclusion, exposure of cells to Cd (up to 75 μM for 24 hours) resulted in inhibitory concentrations 50% (IC50) and 25% (IC25) of 40 and 20 μM, respectively. Real-time PCR analysis of genes associated with epigenetic chromatin modification during early embryonic development indicates that treatment of cells with IC50 and IC25 concentrations results in a greater than 2-fold up-regulation of Aurkcs, Setd1b, and Smyd1. Aurkcs, a member of the Aurora kinase family, has a significant influence on cell proliferation of the pre-implantation blastocyst. Setd1b is a component of the histone methyltransferase complex that trimethylates H3K4 (histone 3, a core histone protein, at position lysine 4), resulting in downstream expression of Oct-4. Smyd1 is involved early in embryonic development of cardiac progenitor cells. The results suggest that Cd up-regulates genes that are important in early mitosis and blastocyst formation, methyltransferase activity, and cellular differentiation, the consequences of which implicate the potential role of the metal to alter the normal processes that are coupled with embryonic development in vivo.

1198 Colony-Forming Cell (CFC) Assays Predict for Increased Clinical Neutropenia Resulting from Combination Therapies.
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Combination therapy can be more efficacious than monotherapy in preventing progression of certain diseases. Unfortunately, this increased therapeutic efficacy often comes with increased hematotoxicity. In a randomized study of 646 patients with Multiple Myeloma, Orlowski et al. (2007) saw clinical neutropenia in 20% of patients receiving Bortezomib alone as compared to 35% of patients receiving Bortezomib and Doxorubicin. Shah et al. (2007) reported clinical neutropenia in
25% of patients receiving Imatinib and Hydroxyurea in patients with recurrent ma- lignant glioma. Some myelotoxic combinations may be unintentional: Hachem et al. (2003) reported myelosuppression in bone marrow transplant patients treated with Linezolid that also happened to be on selective serotonin reuptake inhibitors (SSRIs). To see if in vitro CFC assays could predict the clinical neutropenia ob- served with these combinations, we tested the compounds alone and in combina- tion on CFU-GM inhibition. Normal bone marrow cells and single compounds (range of concentrations) were mixed in methylcellulose-based media (ColonyGel 1102) and plated in 35 mm dishes (3 replicates/concentration). CFU-GM were enumerated on day 14. Compound IC50 values were: Bortezomib (B) 12±μM; Doxorubicin (D) 28 μM; Imatinib (I) 2.6 μM; Hydroxyurea (H) 31 μM; Linezolid (L) 109 μM; and the SSRI Fluoxetine (F) 27 μM.

Drug pairs (B+D; I+H; and L+F) were then tested using a matrix design. When compounds were combined at their IC50 equivalent values, there was an additional inhibition of CFU-GM: 69% with B+D, 84% with I+H and 75% with F+L. These data suggest that the CFU-GM assay may be a useful tool to evaluate drug combi- nations and predict for clinical neutropenia in a variety of circumstances: when concern of increased hemotoxicity with a strategic therapeutic combination exists; when there is limited information on the drug combinations to be employed; when a drug is to be used on pre-treated patient populations. CFC assays may also be use- ful in evaluating compounds with potential myeloprotective effects in combination with myelotoxic compounds.

1199 The Effect of TCDD on the Development of Human Hematopoietic Stem Cells to Lineage-Committed B Cells.

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2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD) is an ubiquitous environmental contaminant that exhibits a broad spectrum of toxicity. Epidemiological studies have identified an association between exposure to TCDD and B cell disorders, such as suppression of humoral immunity and increased incidence of B cell leukemia. Although extensive studies have been dedicated to elucidate the mecha- nism underlying suppression of mature B cell function by TCDD, little is known whether TCDD will alter the sequential progression of human hematopoietic stem cells (HSC) to lineage-committed B cells. To address this question, we established a co-culture system using irradiated human marrow stromal cells as a supportive mi- croenvironment to drive the differentiation of human cord blood CD34+ HSC in the presence of cytokines (IL-3, IL-7, SCF and Flt3L). In three weeks of culture, we observed increased expression of IL-7R, which is a hallmark of lymphopoiesis. Furthermore, generation of lineage-committed B cells was detected by the co- expression of B220 and CD19 in 20% of the cell population. To test the effect of TCDD on B cell development, we treated CD34+ HSC with 1nM, 10nM and 30nM TCDD. Compared to the vehicle control, the proliferation was remark- ably arrested by TCDD in a concentration-dependent manner. We also observed a significant decrease of IL-7R expression in TCDD-treated cells. These data illus- trate establishment of an in vitro model system to drive the development of cord blood HSC to lineage-committed B cells. Preliminary studies showing down regulat- ion of IL-7R suggest a role by TCDD in altering early B cell development.

1200 New Methodology for the Production of Thin-Layered Cardiac Tissue from Human-Induced Pluripotent Stem Cells (hiPSCs) Derived Cardiomyocytes for Use in High-Throughput Electrophysiological Platforms and High-Content Imaging Systems.


We have been developing functional cardiomyocytes derived from human iPSCs for many years. We have previously reported on cryopreserved cardiomyocytes, ReproCardio 2, derived from human iPSCs that demonstrate stable cardiac char- acteristics such as cTnHc, cTnT, MLC-2a, MLC-2v, and CsA43. Using ReproCardio 2 clumps on electrophysiological assay systems, we have further devel- oped our QTempo® assay (on the MEA platform) to better predict the drug in- duced QT interval prolongation (DIQTP).

The use in cardio-toxicity assays requires the reliable assay platform like tissue. We have been developing this technology to make functional thin layered cardiac tis- sue, which has improved beating synchronization and stability; and better reliability for use in cardio toxicity assays. The thin-layer tissue derived from ReproCardio 2 stained positive for cTnT especially around the synchronized beating cell area. We have developed a calcium flux assay (inotropic assay) using the thin layered car- diomyocytes and demonstrated the synchronized myocardial (inotropy) and high content imaging technologies. Furthermore, the thin-layer showed advantages over monolayers, especially its robust signal with changes in serum concentration, which is preferable for high through put screening (HTS) platforms. When using electrophysiology platforms, ReproCardio 2 thin-layer also demonstrated a more robust electrophysiological output similar to that seen with beating cardiomyocyte clumps especially the dose response to DIQTP causing compounds such as E- 4031 and Cisapride. In fact, the results were similar to that obtained in the clinical. The thin-layer cardiomyocytes have shown to be the preferred format giving rise to a robust and sustainable platform for measuring prolonged QT-interval and other cardio toxic parameters such as calcium flux, and in turn will produce a more pre- dictive format for toxicity assays in drug discovery.

1201 Proteomic Analyses of Human Primary Hepatocytes and Stem Cell-Derived Hepatocytes Identifies Hepatocyte Maturity Markers and Shows Improved Phenotype Maintenance in an Air-Liquid Interface Culture Model.

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Failure to predict hepatotoxic drugs in pre-clinical testing makes it imperative to develop better liver models with a stable cultured phenotype. Stem cell-derived models offer promise with hepatocyte-like cells currently considered to be ‘fetal- like’ in their maturity. This is based on limited biomarkers and lacks the required proteomic datasets that directly compares fetal and adult hepatocytes. Here, we quantitatively compare the proteomes of human fetal liver, adult hepatocytes, HepG2 cells and ES-derived hepatocyte-like cells. In addition, we investigate the changes in human fetal and adult hepatocytes when cultured in an air-liquid inter- face format and in extracellular matrix sandwich culture. The proteomes were qual- itatively similar across all samples but hierarchical clustering showed that each sam- ple type had a distinct quantitative profile. Hepatocytes cultured at the air-liquid interface retained a more fetal-like phenotype clumped to fresh cells than (human) conventional culture, which acquired myofibroblast features. Principal component analysis ex- tended these findings and identified sets of proteins as specialized indicators of he- patocyte differentiation. Our quantitative datasets are the first that directly com- pare multiple human liver cells, define a model for enhanced maintenance of hepatocytes in culture and provide new protein markers for determining human hepatocyte phenotype and maturity.

1202 Scalable, HTS/HCI-Amenable and NURR1-Expressing Dopaminergic Progenitor Cells Derived from Human Embryonic Stem Cells.

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One of the most common neurodegenerative diseases, Parkinson’s disease (PD) af- fects approximately 1-2% of the population over age 65. The lack of an appropriate and scalable cell type hinders progress in PD drug discovery and elucidating disease mechanisms. Our goal was to generate human dopaminergic progenitor cells, de- rived from human embryonic stem cells (hESCs), which are homogeneous, ex- pandable for high throughput screening (HTS) and high content imaging (HCI) assays and demonstrate a propensity for dopaminergic differentiation. We isolated cells exhibiting > 95% immunoreactivity for nuclear receptor related 1 protein (NURR1 or NR4A2), a transcription factor essential for generating midbrain dopaminergic neurons. We refer to these cells as DopaPro™ cells, which display normal karyotype, form adherent monolayers and proliferate over multiple pas- sages. DopaPro™ cells were differentiated with leukemia inhibitory factor (LIF), brain derived neurotrophic factor (BDNF), glial derived neurotrophic factor (GDNF), transforming growth factor beta 3 (TGF-β3), forskolin and ascorbic acid (AA). By 3 weeks, differentiated DopaPro™ cells showed tyrosine hydroxylase (TH) expression. We asked whether DopaPro™ cells, as putative midbrain pro- genitor cells, required regionalization signals such as Sonic Hedgehog (SHH) or fibroblast growth factor 8 (FGF8), which are normally used in directed differentia- tion of hESCs to dopaminergic neurons. DopaPro™ cells did not require, nor was their differentiation to TH positive populations significantly enhanced, when cultures were exposed to SHH and FGF8 for 2 weeks prior to LIF, BDNF, GDNF, ...
Growth factors. Results show DopaPro™ cells are a unique, renewable and scalable cell source for PD research.

1203 Conditions for Isolation and Maintenance of Rat Hepatic Stem/Progenitor Cell Colonies.

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Hepatic stem cells, hepatoblasts, and committed hepatocytic and biliary progenitors constitute a stepwise maturational lineage which produces all mature cells of the liver. Hepatic stem/progenitors are localized to the Canals of Hering: peripostral ductular structures derived from ductal plates in fetal and neonatal livers. These peripostral cell niches are enriched in a variety of extracellular matrix molecules including certain types of collagens (III, IV and V) and hyaluronans, non-sulfated polymers of D-glucuronic acid + D-acetylgalactosamine. We have compared the ability of hyaluronans, as well as type III and type IV collagens, to support the proliferation of hepatic stem/progenitors isolated from neonatal and adult rat livers. Cells were isolated either by suspension digestion (neonates) or perfusion digestion with collagenase (adult) and cultured in a serum-free medium (Kubota’s Medium) designed to culture-select endodermal stem/progenitors. Cells from either age seeded onto collagen III or IV (5 ug/cm2) yielded hepatic stem/progenitor cell colonies which expanded for 5-7 days of culture, at which time the mesenchymal cell components overgrew the plating surface. By contrast, a substratum of hyaluronans (5 – 100 ug/cm2), in combination with Kubota’s Medium, promoted coordinated attachment and growth of discrete hepatic stem/progenitors and their mesenchymal progenitor partners. Colonies consisted of hepatic stem/progenitors growing on top of mesenchymal cell progenitors. Neonatal liver-derived colonies had large expansion potential, increasing in area by more than 100-fold by 10 days in culture. It is hypothesized that the expansion potential of adult-derived cells was more limited due to the need for an alternate method of isolation or addition of exogenous growth factors. This novel culture system provides an opportunity to study the effects of hepatotoxins on early lineage stages consisting of rat hepatic stem/progenitor cells.

1204 The In Vitro Human CFU Assay Combined with Automated Image Analysis of Colony Morphology Markedly Improves Assessment of Hematopoietic Toxicity of AZT and Other Nucleoside Inhibitors.

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Antiviral nucleoside analogs target viral polymerases but can also inhibit cellular polymerases, leading to a decrease in cell proliferation and ultimately suppression of hematopoiesis, resulting in anemia and neutropenia. For assessment of hematotoxicity in this compound class, the FDA’s Guidance for Industry: Antiviral Product Development—Conducting and Submitting Virology Studies to the Agency recommends the use of the colony-forming unit (CFU) assay, which utilizes primary hematopoietic stem and progenitor cells. In 2003, the European Centre for the Validation of Alternative Methods (ECVAM) published a standardized CFU assay methodology that is validated for the prediction of the maximum tolerated dose (MTD) of myelosuppressive compounds (Pessina et al., Tox Sci. 2003, 75:355-367). This in vitro assay is a key tool for assessing toxicity prior to initiation of costly in vivo studies and human clinical trials. Unfortunately, in these standardized CFU assays, nucleoside inhibitors such as azidothymidine (AZT) do not exhibit high toxicity (IC50 = 100-200 µM), despite the fact that they are well known to perturb hematopoiesis in patients. Although AZT does not dramatically reduce the number of hematopoietic colonies in the CFU assay, does it does have a dramatic effect on their morphology. To quantitate changes in colony morphology, we utilized the STEMvision® platform for automated advanced image analysis in conjunction with the CFU assay. Our results indicate that IC50 values for AZT based on the quantitation of morphological changes such as colony size were ~20-fold lower (5-10 µM) than the IC50 based on colony numbers (100-200 µM). These data suggest that quantitation of colony morphology is a key parameter to predict drug potency and in vivo toxicity in this drug class.

1205 Investigation of a Redox-Sensitive Predictive Model of Mouse Embryonic Stem Cell Differentiation via Quantitative Nuclear Effect Protection Assays and Glutathione Redox Status.

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Mouse embryonic stem cells (mESCs) recapitulate developmental signals that occur in vivo and are amenable to high-throughput profiling of chemical-induced effects. In silico models of impaired mESC differentiation identified 19 ToxCast™ assays that distinguished chemical effects on cardiomyocyte differentiation versus cytotoxicity (Wilcoxon rank sum, p<0.03). Taken together, these assays connect a redox-sensitive pathway(s) into a likely target of a chemical set that impaired mESC differentiation. To evaluate this predictive model, a custom quantitative nucleae protection assay (qNPA) array (HTG Molecular Diagnostics) of 41 redox-sensitive targets and differentiation markers was designed. The concentration-dependent effects of twelve ToxCast chemicals were evaluated using the qNPA array. The highest concentration produced AC 20 cytotoxicity in mESCs and the remaining were half logarithmic decrements. Cells were harvested on days 4 and 9 of culture. The prediction model correctly identified that pyridaben would disrupt redox signaling in mESCs. Expression of the redox-responsive transcription factors Nrf2, Hif1A and Mtf2 increased upon exposure. Pyridaben showed significant concentration-dependent effects on markers for endoderm, ectoderm and mesoderm differentiation on day 4 of culture (decreased Fgf5, Otx2, and Fgf8 expression and increased Tbx3 expression) without producing a 50% decrease in cardiomyogenesis in the standard assay on day 9. Preliminary data evaluated the redox status (glutathione/glutathione disulfide) of mESCs at 3, 6, 9 and 24 hours following chemical exposure and indicated a unique pattern of redox status for predicted redox disrupting chemicals. These experiments document the importance of using multiple differentiation endpoints and support the linkage of our predictive model to altered differentiation in chemical profiling. This abstract does not reflect EPA policy.

1206 Arsenic Trioxide Alters Regulation of Differentiation in Mouse Embryonic Stem Cells.

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Arsenic trioxide (As2O3) is a known human carcinogen yet it is currently approved by the Food and Drug Administration as a chemotherapeutic agent for the treatment of acute promyelocytic leukemia. As2O3 alters cellular differentiation by mechanisms of which are yet not clearly understood. Thus, in the current study, we investigated the influence of As2O3 on differentiation of mouse embryonic stem cells (mESCs). These pluripotent stem cells are derived from the inner cell mass of the blastocyst, and are capable of continuous proliferation, self-renewal and differentiation into the three germ layers. Undifferentiated cells express early specific stem cell markers, including Oct-3/4, Sox-2, and Nanog. Sox-2, which have an important role in developmental regulation and differentiation. In our initial studies, 1.0 µM to 6 µM As2O3 reduced cell viability over 24-hour with increasing concentrations, whereas significant reductions were observed at or below 1 µM only with 48-hour exposures, a concentration which is equivalent to therapeutic administration. End point PCR analysis revealed that increasing concentrations of As2O3 reduced gene expression of stem cell markers. This supports the western blot results, protein levels decreased with decreased in gene expression. Western blot analysis for Oct-4/3 and Sox-2 protein expression levels are crucial for maintaining mESC in their undifferentiated state, the reductions in the presence of arsenic imply the loss of “stemness”. Consequently, it appears that As2O3 exposure reduces the ability of the stem cells to maintain proper state of differentiation.

1207 Vascular Endothelial Cells Exposed to PCB 153 Show Increased Expression of Stem Cell Markers.


The contribution of PCB exposure to the development of vascular disease is an area of research that has received little attention. The stability of these chemicals has allowed them to persist in the environment and consequently continues to expose the general population to PCBs. Since PCBs have been reported in human blood, vascular toxicity from PCB exposure is a public health concern. Vascular lesions from the lungs of patients with severe PAH have been characterized with excessive EC proliferation and markers of angiogenesis. The exact cell type from where vascular
lesions originate from is not clear. However, our studies have focused on ECs as the origin of vascular injury because EC stem cells have been reported to be involved in vascular injury. The theory that malignancies depend on stem-like cells for proliferation has received much attention, but there have been few studies which support a pathogenic role for stem cells in vascular disease. Thus, the objective of this study was to determine whether exposure to the PCB congener 153 can increase the level of stem cell markers in vascular ECs. ECs overexpressing ID3 were used to study the effects of PCB153 in 3D spheroid cultures. Co-culture model of EC with SMGs and fibroblasts was used to study the expression of stem cell markers in a monolayer by confocal microscopy. The effect that PCB153 had on cell cycle progression was also determined by flow cytometry. We observed PCB153 significantly increased expression of stem cell markers CD34+, CD133+, VEGFR-3 in spheroids overexpressing ID3. ECs exposed to PCB153 showed stable spheroid formation in B27 medium at 10 days. ECs exposed to PCB153 in co-cultures showed increased expression of stem markers CD34+ and Nanog as well as increased angiogenic phenotype. Cell cycle analysis confirmed that more stem-like cells exposed to PCB153 were in G1 phase. Our results show that PCB153 exposure increased expression of stem cell markers and may be related to mechanisms by which vascular disease depends on stem-like cells for proliferation.

**1208 Evaluation of Ionotrophic and Chronotrophic Compounds In Vitro Using Human iPS-Derived Cardiomyocytes and Impedance-Based Contractility Assay.**

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The β-adrenergic receptor (βAR) signaling system is one of the most powerful regulators of cardiac ionotrophic and chronotropic function, βAR antagonists, commonly known as β-blockers (β-blockers) have been used for decades to treat hypertensive, ischemic heart disease, some arrhythmias, and congestive heart failure. Up to date, 3 types of β-blocker drugs have been developed for therapeutic purpose. In order to better understand the cardiac effects of different types of β-blockers, we have developed an in vitro assay using human iPS-derived cardiomyocytes in conjunction with impedance measurement. We evaluated the activities of six β-blocker drugs and compounds including (1) non-selective β-blockers; (2) cardioselective β-blockers; and (3) nonselective α and β-blockers. The data as measured by impedance readout revealed that the β-blockers from different classes displayed subtle but unique profile of acute impedance changes post compound addition. With the exception of atenolol and metoprolol, pindolol, alprenolol, propranolol and carvedilol significantly decreased the cell beating rate (BR) and even temporally led to beating arrest at the highest tested concentration (5 μM). In addition, the 1 hour pretreatment with all these four compounds at higher doses abolished the isoproterenol-induced positive chronotropic effects on BR. Carvedilol was the only drug that appeared to profoundly reduce cell beating amplitude (CA) in a dose-dependent manner. The in vitro assay used here indicates that human iPS-derived cardiomyocytes respond similarly to the β-blockers from the same group. However, each type of β-blockers generates distinct profiles and do have different chronotropic and ionotropic effects which we will summarize in the current poster.

**1209 Selection of CYP3A4 hESC-Derived Hepatocytes for Drug Metabolism and Toxicity Assays.**

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The ability to produce human hepatocytes that express adult levels of drug metabolizing enzymes including CYP3A4 in a stable and reproducible system will greatly benefit the drug development process by decreasing costs, reducing animal studies, and increasing drug safety. Many groups are trying to develop hepatocytes from human stem cells to address the lack of availability, quality, and reproducibility of primary hepatocytes, however, most protocols produce immature hepatocytes with minimal expression of the adult enzymes related to drug metabolism. Since CYP3A4 is the most abundant P450 expressed in adult liver and metabolizes 50% of drugs in the market, it is imperative that this system expresses this enzyme. To this end, we previously reported the generation of a hESC reporter line (3A4BLA) containing humanized beta-lactamase (hBLA) inserted in frame with the CYP3A4 gene. Here we show that 3A4BLA-hepatocytes can be used to: 1) monitor the differentiation of mature hepatocytes; 2) sort for mature hepatocytes; 3) monitor drug induction of CYP3A4; and 4) develop in vitro assays for drug metabolism and toxicity. Using an optimized protocol for hESC-hepatocyte differentiation, we demonstrate CYP3A4 in 25-40% of the cells and show that these cells produce levels of CYP3A4 mRNA approaching primary hepatocytes, express high levels of ALB, and store glycogen. More importantly, they metabolize testosterone and midazolam and respond properly to inducers of CYP3A4 like Rifampicin and Phenobarbital. With this system we can obtain a relative measure of CYP3A4 expression on a per cell basis in response to compounds and correlate those data with cell metabolism, CYP3A4 activity, and secretion of albumin and urea. Taken together, these data demonstrate development of a robust system with the potential of purifying human hepatocytes with more mature functions, as well as assaying drug effects on CYP3A4 expression, activity, and metabolism.

**1210 Adapting the MTT Assay for Use with Human Embryonic Stem Cells (hESC).**

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hESC are difficult to adapt to 96-well plate assays because they survive best when plated as colonies, which are difficult to count and plate accurately. To address this problem, two protocols were developed to perform the MTT assay using hESC. In the first protocol, plating was done with Rho-associated kinase inhibitor (ROCKi), which allows accurate counting and plating of single hESC. The second protocol involved plating hESC as small colonies without ROCKi but with adaptations to allow accurate counting and plating of small colonies. In the ROCKi protocol, 5,000 cells/well are counted using a hemocytometer then plated and incubated for 96-hours, while the small colony method calls for 20,000–30,000 cells/well followed by 72-hours of incubation. To enable rapid and uniform plating of small hESC colonies, we developed a spectrophotometric method to accurately determine hESC concentration. The percent transmittance for multiple readings of the same sample produced a coefficient of variation (CV) of 1.5%. Next, we determined the cell concentration needed to carry out an experiment in a 96-well plate by plating various cell concentrations and determining the optimal concentration needed for a 72-hour experiment. Finally, to confirm that the correct cell number was pipetted accurately into each well, control wells were subjected to the MTT assay and used to calculate the COV, which ranged from 0.7 to 8.5%, showing that pipetting of small colonies was uniform. The small colony protocol was validated using the NIH Plate Uniformity and Signal Variability Assessment, which tested for signal separation, drift and edge effects. When comparing the ROCKi protocol to the small colony protocol, it was found ROCKi caused a shift in the IC50 from 1.34e-3 to 1.45e-4 M. In addition, hESC morphology was altered with ROCKi treatment, which appeared to stress the cells, while cells in the small colony protocol appeared healthier, tightly packed, and cobblestone-like. Both protocols allow the MTT assay to be carried out rapidly and accurately with high reproducibility between replicate experiments using hESC.

**1211 Differential Regulation of Pluripotency Maintenance and Differentiation Genes in Undifferentiated Human Embryonic Stem Cells (hESC) Exposed to Methyl Parathion (MP) and Its Active Metabolite Methyl Paraoxon (MPO).**


Given their widespread use, ease of procurement, and potential for serious health consequences if deployed by terrorists, toxic industrial chemicals (TICs) represent a real threat to warfighters and civilians at home and abroad. Unfortunately, for a vast number of TICs, including the widely-used organophosphate insecticide, MP, there is incomplete knowledge regarding the basic molecular consequences of exposure in humans. Although the literature suggests diverse toxicological consequences for MP exposure during early fetal development, it is primarily based on human epidemiological studies, in vivo animal studies, or in vitro studies using immortalized, transformed cell lines. Furthermore, the relative contribution and the molecular mechanism(s) of action of the parent compound, MP, versus the active metabolite, MPO, has not yet been elucidated in an early developmental system. In an in vivo model, the MP is converted by hepatic and extra hepatic phase I and II metabolism to MPO, and the relative ratios can vary depending on the in vivo system. Given that undifferentiated hESCs do not produce the enzymes necessary to carry out MP metabolism very early human development.

In this study we compared the expression of 84 genes known to be important in the maintenance of pluripotency and differentiation in hESCs cells following a 24 hour exposure of cells to either MP or MPO. These RT-PCR data indicate several key differences in the expression of genes following exposure to MP and MPO. The results of this study have opened a new avenue toward understanding how MP and its active metabolite MPO specifically interfere with some of the earliest steps in human embryonic development.
In this study, we used a "toxicoproteomics" approach and SILAC (Stable Isotope Labeling with Amino acids in Cell culture) to study the changes in the proteome of the human hepatocellular carcinoma cell line HepG2 following treatment with a set of seven compounds (Labelatal, Nicotinic Acid, Diclofenac, Amodexicilllin, Cefaclor, Rifampin and Tacrine) previously reported to be involved in DILI. We identified over 4,000 proteins by high-resolution mass spectrometry with a false discovery rate of 1%. We used MaxQuant software for protein identification and the quantitative analysis of data from mass spectrometry and DAVID Bioinformatics - GO (gene ontology) analysis tools to determine protein changes and biological processes/pathways affected by the treatment of cells with the drugs. Our results indicate that the levels of some enzymes with potential effects in drug toxicity were reduced when treated with the compounds. These include carnitine O-acyltransferase activity (involved in the metabolism of fatty acids), nucleotide excision repair (DNA repair function) and Glutathione S-transferase A1 (detoxification of electrophilic compounds). Biological processes generally affected as determined by the GO analysis include the response to metal ion and nitrogen compound biosynthetic processes. These results provide further insight into possible cellular mechanisms linked to the toxic activities of the drugs analysed. Undoubtedly, quantitative proteomics could play a pivotal role in deciphering toxicity mechanisms at the molecular level.
Our analysis detected and corrected for small but persistent batch effects related to sample processing and employed multi-dimensional methods for differential statistics in >1000 samples. Each transcriptomic drug response profile differentiates drug-specific on- and off-target activities between cell types, within drug classes, and between drug classes, as well as estimating relative molecular pathway activity between drugs within a class. For example, preliminary analysis of TZD-class re- sponses ranked the on-target “insulin receptor signaling pathway” for pioglitazone at 6th, but rosiglitazone at 159th, and troglitazone at 259th, while off-target “can- cer” and “cell remodeling” pathways dominated the top 10 signaling pathways for troglitazone and rosiglitazone, consistent with clinical history of these drugs. CONCLUSIONS: Methods combining human-relevant systems with mining drug-specific profiles in conjunction with external drug information databases (FDA’s AERS, Altman’s OFFSIDES, EPAs ToxCast, Broad’s Connectivity Map) serves to enhance ability to make broader human vascular drug response predic- tions.

1217 Towards a Platform of BAC-GFP Transgenomics-Based Pathway of Toxicity Reporters for Automated High-Content Imaging-Based Chemical Safety Assessment.


Adaptive cellular stress responses are paramount in the healthy control of cell and tissue homeostasis after cell injury during hypoxia, oxidative stress or unanticipated side-effect of medications and other chemical exposures. Genome-wide transcriptomes analysis has revealed the detailed cellular stress response landscape and thereby the diversities of organelles and cell functions that are being affected upon exposure to diverse chemicals. Individual toxicological relevant stress responses have also been termed pathways of toxicity (PoT). We find that activation of PoT occur well before the typical ultimate outcome of chemical cell injury: cell death by necrosis or apoptosis. Understanding the activation of PoT caused by chemicals is complex because many simultaneous biochemical cellular perturbations may occur thereby affecting different PoT in parallel or a defined order. To increase our understanding of chemically-induced PoT activation and its contribution to safety assessment we believe that a time-resolved, sensitive and multiplex readout of chemical-induced toxicological relevant PoT will be essential.

For that purpose, we are currently developing a automated live-cell high-content imaging-based platform containing a panel of distinct PoT reporter cell lines. To conserve the endogenous regulation we tag selected target genes with GFP using BAC-transgene-omics approaches. So far we have generated 23 different human he- patoma HepG2 reporter cell lines that define autophagy, mitochondrial organization, Golgi structure, nucleosomes, ER morphology and stress, oxidative stress re- sponse and DNA damage responses.

Here we report the current status of our BAC-GFP PoT reporter platform and demonstrate that individual reporter cell lines are sensitive to their corresponding model stress responders. We anticipate that ultimately a phenotypic PoT profiling platform will allow a high throughput and time-resolved classification of chemical-induced stress responses in the safety assessment of chemicals.

1219 Nonclinical Toxicology Profiling of the First-In-Class Sodium-Glucose Co-Transporter 2 (SGLT2) Inhibitor Dapagliflozin.


Dapagliflozin (Dapa) is a novel SGLT2 inhibitor that directly increases urinary glu- cose excretion as a mechanism for treating diabetes. To support its safe use during clinical development, Dapa was evaluated in a rigorous battery of in vitro and in vivo nonclinical safety studies. In vitro screening of >300 diverse targets indicated no significant off-target activities for Dapa or its primary metabolite. The complete set of in vivo safety studies included dosing up to 12-months in dogs and a lifetime (~2 years) in mice and rats. As expected, most of the effects were related to Dapa-induced pharmacology (i.e. SGLT2 inhibition) such as glucosuria with osmotic diuresis and mild loss of electrolytes, and limited body-weight gains despite increased food consumption. At extremely high doses, Dapa also caused intestinal glucose malabsorption. This effect only occurred in rats at extremely high doses (>2000x clinical exposures) and was related to increased intestinal calcium absorption, a consequence of exacerbated carbohydrate fermentation in the rodent gut related to SGLT1 inhibition, which ultimately contributed to thickened trabeculae in long bones and soft tissue mineralization. Of particular importance, chronic Dapa treat- ment at >600x clinical exposures did not result in hyperplasia or any adverse effect on the kidneys or urinary tract nor at > 100x clinical exposures any increases in the incidence of or shortening in the latency period for tumor development. There was also no indication that SGLT2/- mice had adverse urinary tract morphology or function when compared to their wild type counterparts in a 15-month observa- tional study. Thus, the pharmacologic activity of Dapa in nonclinical species and the rigorous attempts to test supra- and off-target pharmacology allowed us to throroughly evaluate its potential toxicity profile, providing us with confidence in its safety in patients with diabetes.

1220 Bone Assessments in Enhanced Pre-Postnatal Toxicity Study in Pregnant Cynomolgus Monkeys with up to 6-Months Postnatal Evaluation.

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Assessment of potential test article-related bone liability is becoming increasingly important, not only in the mature skeleton but also in the developing skeleton. Skeletal assessments were developed to evaluate potential liability in pregnant mon-keys and offspring in an enhanced prenatal to postnatal development (ePND) toxicity study. Assessment in mothers during gestation and lactation included bone markers C-telopeptide (CTXs), tartrate-resistant acid phosphatase active isofrom 5b (TRAP5b), osteocalcin (OC), and bone specific alkaline phosphatase (bALP). In infants, assessment included bone markers, radiology, ex vivo bone densitometry (μCT) and biomechanical testing.

Control mothers had generally high values of bone markers during early pregnancy and lactation. At GD20, most had OC levels above the upper limit of quantifica- tion then decreased at other time points during gestation, and increased during lac- tation. bALP and CTx increased at the end of the gestation and during lactation. TRAP-5b increased during pregnancy but decreased during lactation. In control in- fants, OC, bALP and TRAP-5b were increased up to 2, 4 or 10 fold, respectively, compared to historical data for 3-4 year-old animals. Radiographs were evaluated for skeletal abnormalities and bone length: age-related changes in ephiphysial mineralization, growth plate morphology, and increases in bone length were noted over time. Femur and lumbar vertebrae geometry (μQCT) changed with increasing age with lower increases in density. Femur 3-point bending and vertebral compression showed increases in strength parameters with age (2-fold between BD1 and BD 180).

Skeletal changes were consistent with gestation and lactation in mothers, and with development and growth in infants. These data support the use of bone markers, densitometry, radiology, and biomechanical testing in toxology studies, including ePND studies, to provide a comprehensive evaluation of the maternal and develop- oping skeleton in nonclinical drug development.
Characterization of Batracylin-Induced Renal and Bladder Toxicity in Rats.

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Batracylin (NSC-320846; BAT) is an investigational anticancer agent that reached Phase 1 clinical trials. BAT is a dual inhibitor of DNA topoisomerases I and II and induces histone gamma-H2AX, a biomarker of DNA damage in vitro. Hermorrhagic cystitis was one of the dose limiting toxicities observed during clinical trials, and bladder and kidney were postulated to be responsible for the hematurcia observed in the clinic. We investigated the mechanism of bladder and renal toxicity in Fisher 344 rats, a physiologically relevant model. The studies were designed to 1) examine the effect of BAT administration on rat bladder histology, 2) further characterize the previously reported renal toxicity of BAT, and 3) measure DNA damage in the kidney and bone marrow after BAT administration in rats using gamma-H2AX immunofluorescence. Once daily oral administration of 16 or 32 mg/kg BAT to Fisher 344 rats for 4 days caused overt toxicity. Abnormal clinical observations, adverse effects in clinical pathology, urinalysis, kidney and bladder were seen. Gamma-H2AX immunofluorescence indicated DNA damage in kidney and bone marrow. Furthermore, after administration of BAT, defects in the superficial and intermediate urothelial layers were observed using scanning electron microscopy. The maximum tolerated dose is estimated to be <16 mg/kg/day. MesnExTM is known to reduce the incidence of hemorrhagic cystitis induced by ifosfamide or cyclophosphamide, but BAT toxicity in rats was not alleviated by twice daily intraperitoneal administrations of 80 mg/kg mesna. Thus, the mechanism of BAT-induced bladder and renal toxicity was not mediated by urotokinetic mechanisms similar to those of ifosfamide or cyclophosphamide. These studies show that BAT causes renal and urothelial damage in rats that may be related to DNA damage, a previously identified mechanism of action for BAT. Work supported by NCI Contract N01-CM-42203 and N0-CM-2011-00628.

NKTR-181, a Novel Opioid Agonist, Demonstrates Mu-Opioid Agonist Effects and Minimal CNS Side Effects in Both Rat and Dog.


NKTR-181 is a novel orally available mu-opioid agonist designed using Nektar's advanced polymer conjugate technology to achieve slower kinetics of entry into the brain relative to other opioids, while still producing analgesic efficacy. Previous studies of NKTR-181 in rodent models have demonstrated a clear separation of analgesia from unwanted CNS side effects compared with oxycodone. To assess the safety of this novel opioid, NKTR-181 was evaluated after daily oral dosing for 2 and 13 weeks in rats and dogs at doses up to 600 and 75 mg/kg/day, respectively. Clinical observations, body weights, clinical chemistry, and histopathology were performed and NKTR-181 plasma concentrations were analyzed by LC-MS/MS. Findings in both species were consistent with the pharmacology of a mu-opioid agonist; no off-target toxicity was observed. In rats, the major clinical sign was foot sores due to increased paw chewing at 200 to 600 mg/kg/day, a known opioid effect in rodents. Mild to moderate gait and activity changes were noted at 600 mg/kg/day in a neurobehavioral assessment in the 2-week study. The incidence of clinical findings diminished with repeated dosing. Decreased mean body weights relative to controls was observed in males at ≥200 mg/kg/day and in females at 600 mg/kg/day. In dogs, major clinical signs were post-dose emesis at doses ≥15 mg/kg/day and mild, transient hypoactivity at 75 mg/kg/day. Accommodation to both emesis and hypoactivity was observed. Decreased mean body weights relative to controls were observed at 75 mg/kg/day in the 13-week study. No adverse clinical pathology or histopathological changes were noted in the rat or the dog. The NOAEL was determined to be 200 mg/kg in rats and 75 mg/kg in dogs (human equivalent doses of 32 and 42 mg/kg, respectively). Plasma levels (AUC) at the NOAEL were 170-634 and 50-155 ng/mL, in rats and 53 and 326 ng/mL in dogs. These studies demonstrate that NKTR-181 is well tolerated and produces minimal CNS side effects in both rat and dog.

Safety Assessment of NU100 in a 6-Month Cynomolgus Monkey Study.

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NU100 is recombinant human interferon beta-1b (IFN beta-1b) being developed for treating multiple sclerosis. NU100 has a better purity profile (aggregate-free and HAS-free) compared to other marketed products. A GLP monkey study for NU100 safety assessment was conducted which was originally planned as a 6-month subcutaneous injection study with an 8-week recovery. Male and female cynomolgus monkeys were assigned to 5 groups (4 or 6 animals/group), and were dosed once daily or every other day with NU100, Betaseron and placebo at dose levels of 0, 0.01, 0.06, or 0.28 mg/kg/dose. Animals were monitored clinically, and blood samples were collected for evaluation of serum levels of IFN beta-1b, neopterin (a biomarker for IFN beta-1b pharmacodynamics), and anti-drug antibodies (ADA). NU100-related adverse clinical findings were noted during the study. IFN-beta-1b serum concentration peaked in circulation within 2 to 4 hours after administration of NU100 on Days 1 and 15 with a higher level on Day 15; however, by Weeks 18 and 22, IFN beta-1b level returned to the baseline levels. Similarly, neopterin concentrations increased on Day 1 following administration of NU100 in a dose-dependent manner. Mean predose neopterin concentrations remained slightly elevated on Day 15 compared with Day 1 baseline levels. Nevertheless, no further induction of neopterin was noted on Day 15. By Weeks 18 and 22, neopterin concentrations returned to predose levels. At week 2, 12.5% animals treated with NU100 and 37.5% treated with Betaseron were confirmed positive for ADA. All NU100 treated primates were confirmed positive for ADA at weeks 2, 4, 8, 12, and 18 during the study. Due to the loss of IFN-beta-1b activity resulting from the generation of anti-IFN beta-1b antibodies, the dosing phase of the monkey study was terminated one month earlier than planned, following discussion with and approval from the FDA. In conclusion, NU100 was well tolerated by monkeys in this chronic toxicology study.

BMP31510 Is a Novel Regulator of Mitochondrial Function That Mitigates/Rescues Drug-Induced Toxicity—Evidence in Drug-Induced Cardiotoxicity and Cancer Chemotherapy Using In Vitro and In Vivo Models.

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In a large cohort of drugs routinely used in clinical practice, disruption of mitochondrial function represents a common thread associated with incidence of organ specific toxicities. An example is the incidence of cardiotoxicity associated with anti-diabetic drugs, cancer chemotherapeutics and anti-retroviral agents. Ubidecarenone is a mitochondrial resident molecule with multiple functionalities including electron transport and generation of ATP. The delivery of ubidecarenone to the mitochondria is a challenge due to its physiochemical properties. BMP31510 is a nanodispersin formulation containing ubidecarenone incorporated into a novel lipid mixture with improved ability to deliver ubidecarenone with ability to influence mitochondrial function. The effect of BMP31510 on the toxicity profiles of several cancer chemotherapeutic agents was tested in cell cultures and animal models of cancer and on the cardiotoxicity potential of an anti-diabetic drug in primary cultures of normal human cardiomyocytes. BMP31510 significantly enhanced the efficacy of multiple chemotherapeutic agents used for treatment of cancers of the pancreas, prostate, breast and colon in in-vitro and in-vivo models. In addition, BMP31510 mitigated chemotherapy induced toxicity in primary cultures of normal human liver, fibroblast and prostate cells. Furthermore, BMP31510 reversed mitochondrial dysfunction resulting from exposure to an anti-diabetic drug. BMP31510 represents a novel technology effectuating a multimodal physiological response in which it does not interfere with the normal therapeutic mechanism of action of chemotherapy drugs, yet confers protection to normal tissues. BMP31510 is currently in clinical trial as a mono-therapy for solid-tumors with no reported adverse effects.

Etirotinotec Pegol Nonclinical Toxicology Studies Establish a Margin of Safety to Support an Every Three-Week Clinical Dosing Schedule.


Background: Etirotinotec pegol, formerly NKTR-102, is a unique targeted topoisomerase I inhibitor designed for prolonged tumor cell exposure. Etirotinotec pegol demonstrated a favorable tolerability profile in nonclinical and clinical studies with improved safety over irinotecan. Toxicology studies were conducted in dogs to further evaluate etirotinotec pegol safety and establish dose-related toxicities. Methods: Etirotinotec pegol was administered as an IV infusion to dogs for 3 months using a q14d schedule at doses up to 30 mg/kg (600 mg/m2) irinotecan equivalents. Direct comparison to irinotecan was evaluated in dogs using a q7dx4 schedule (Persson et al, 2008). Clinical observations, clinical chemistry, and histopathology were performed and toxicokinetic samples were collected and assayed by LC-MS/MS for etirotinotec pegol, irinotecan, SN38, SN38-Glucuronide,
and APC. Results: Erititnotecan peg-treatment dogs (25 mg/kg, 500 mg/m²) had reduced neuropenia and diarrhea and no mortality compared to irinotecan-treated animals at the same dose when given q4d. When given q1d for 3 months, erititnotecan peg-treatment dogs (30 mg/kg, 600 mg/m²) had no neuropenia and minimal diarrhea at SN38 exposure levels up to 5-fold above those in patients given 145 mg/m² erititnotecan peg on a q2d schedule. Body weight gain suppression and decreased food intake were the primary side effects observed at this erititnotecan peg dose and schedule. No test-article related alopecia was noted in animals treated with erititnotecan peg in any of the toxicology studies in contrast to literature reports of aloepecia in both humans and in animals treated with irinotecan. Conclusions: Nonclinical toxicology studies show that erititnotecan peg produces markedly less neuropenia and diarrhea with no mortality when compared to irinotecan. Erititnotecan peg is well-tolerated for 3 months at doses that provide a good safety margin (based on SN38 exposure) to support a clinical regimen of 145 mg/m² given every 3 weeks.

**1226** Transient Thrombocytopenia without Coagulopathy in Rats following Single IV Bolus of Oxycyte®, a Perfluorocarbon (Pfc) Based Oxygen Carrier.

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Oxycyte®, a 60% w/v perfluoro(1-butylcyclohexane)FrBu) intravenous emulsion, is being developed for treatment of traumatic brain injury (TBI). Transient thrombocytopenia (TTP), a known class effect of PFCs, may pose a risk for TBI patients. A rat model of intracranial hemorrhage (ICH) is under development to ascertain whether TTP subsequent to Oxycyte administration exacerbates ICH and whether platelet transfusion will ameliorate TTP associated ICH. As part of model development, studies were conducted to determine hematology, coagulation, cytokine, and PK profiles following Oxycyte administration. Male rats received a single IV bolus of Oxycyte at 3, 6, or 12 mL/kg. Blood was collected via jugular cannula (n=3/group) at 10 post administration time points and analyzed for FrBu concentration. Additional rats (n=5/group/time point) were sacrificed on Study Days (SD) 2, 3, 4, 5, 7, 10, 14, and 30 for hematology and blood cytokine analyses. Blood T1/2 following 3, 6, or 12 mL/kg was 4.2, 3.9, and 6.3 hours respectively. AUC was 12- and 48-fold higher for rats receiving 6 and 12 mL/kg compared to 3 mL/kg. There was a 30-40% decrease in PLT counts on SD3-5 in rats receiving 3 or 6 but not 12 mL/kg suggesting Oxycyte-induced PLT sequestration (TTP) rather than PLT destruction. Mean platelet volumes were increased in all Oxycyte groups through SD7 with a decrease in reticulocyte (RTC) counts and elevated fibrinogen levels through SD5. Except for elevated fibrinogen concentrations, there were no significant changes in coagulation parameters. No elevations in IL-6, TNF-α, and IL-1β levels were detected. Liver and spleen weights were elevated at all time points. RTC changes were considered a pharmacologic response to Oxycyte with no concurrent changes in erythrocyte counts. Increased liver and spleen weights were expected effects due to removal of Oxycyte by the reticuloendothelial system, the known mechanism for clearing PFC particles.

**1227** Immunomodulatory Activity of Orphan Drug Elmiron® in Female B6C3F1/N Mice.

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Intestinal cystitis/painful bladder syndrome (IC/PBS) is a chronic disorder characterized by bladder discomfort and urinary urgency in absence of identifiable infection. Elmiron® (EMR; sodium pentosan polysulfate) is the only approved oral therapy for treatment of IC/PBS. Based on the lack of chronic toxicity data and potential for long term use in the treatment of IC/PBS, NTP conducted 90-day and 2-year toxicity studies for EMR. The results suggested that EMR could potentially modulate the immune system. Therefore, this study was conducted to evaluate the immunomodulatory effects of EMR when administered for 28 days via oral gavage to female B6C3F1/N mice, at doses of 63, 125, 250, 500 and 1000 mg/kg. Mice treated with EMR had a significant increase in absolute liver weights (500 and 1000 mg/kg) and natural killer (NK) cells (250 and 1000 mg/kg) were significantly increased. EMR treatment did not affect the humoral immune response (antigen specific antibody response to sheep red blood cells [SRBC] or cell-mediated immunity (mixed leukocyte response). However, innate immune responses such as phagocytosis of radiolabelled SRBC by liver macrophages (1000 mg/kg) and NK cell activity were enhanced (500 and 1000 mg/kg). Further analysis using a disease resistance model showed that EMR-treated mice demonstrated significantly increased anti-tumor activity against B16F10 melanoma cells at the 500 and 1000 mg/kg doses, where the numbers of tumor nodules were decreased by 55% and 68%, respectively. Collectively, we conclude that EMR administration stimulates the immune system, increasing numbers of specific cell populations and enhancing phagocytosis and NK cell activity in female B6C3F1/N mice.

**1228** Assessment of Long-Term Preclinical Safety of Inhaled Technosphere® Particles and Afrezza® Inhalation Powder.

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The inhalable insulin, AFREZZA® inhalation powder and novel excipient, FDLP (fumaryl diketopiperazine which self assembles into Technosphere® Particles), were evaluated in nonclinical safety studies for the treatment of diabetes. Daily doses of either Technosphere particles or AFREZZA inhalation powder were administered by nose-only route in chronic repeat dose and carcinogenicity studies in rats or oronasal route in dogs. With the exception of transient exaggerated pharmacological effects of insulin, inhalation administration for up to 104 weeks in rats and 39 weeks in dogs was well-tolerated. There were no adverse effects on body weight, food consumption, ophthalmoscopy, clinical pathology parameters or electrocardiography (dogs). There were no indications of carcinogenic potential or pro- liferation related to AFREZZA inhalation powder in pulmonary tissues by immunostaining using proliferating cell nuclear antigen assay. Non-proliferative lesions were limited to minimal goblet cell hyperplasia and eosinophilic accumulation in the olfactory/respiratory epithelium in rats; and minimal neutrophil infiltration in lung in dogs. Maximum systemic concentrations of insulin and FDLP were rapidly attained. Terminal half life was approximately 20 and 60 min for insulin and FDLP, respectively. In dogs, systemic exposure of FDLP and insulin was generally dose proportional with no sex differences or accumulation. In rats, insulin exposures were generally higher in females compared to males. Exposure to insulin was related to doses received in the stomach and the increased glucose levels. Based on the data from these chronic inhalation studies, the no adverse effect level was established as the highest studied doses for AFREZZA inhalation powder and Technosphere particles in both species.

**1229** NF-κB Activation in the Hippocampus during Multiple Subthreshold Exposures to Seizurogenic Compounds.

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Drug-induced seizures have been documented for broad classes of pharmaceuticals including CNS and Non-CNS targeted drugs. A better understanding of the early molecular signaling events involved in promoting seizures is necessary to identify potential proconvulsant liability of new pharmacologic agents earlier in the development process. The NF-κB pathway is involved in regulating a number of stress genes and its activation may conceivably be an early indicator of potential seizure liability. Presently, we employed a NF-κB-dependent GFP reporter mouse to investigate the role of NF-κB signaling in rendering hippocampal neurons hyper-excitability. Utilizing EEG recordings and video documentation we established a sub-threshold dose level of the seizurogenic compound kainic acid (KA). Transgenic reporter mice were exposed to multiple sub-threshold doses of KA and regional and cell specific NF-κB activity was assessed after each dose. Under control levels reporter expression was absent in the hippocampus except for a slight basal expression in the CA3 pyramidal layer. Upon multiple exposures to KA, a pronounced expression of the GFP reporter was observed in the stratum molecular, dentate gyrus molecular layer and in the dentate hilus. Additionally we exposed reporter mice to multiple low levels of a different seizurogenic compound, pentylentetrazole (PTZ). After multiple doses of PTZ, a global increase in GFP expression was observed. Comparison of the two compounds suggests a regionally selective expression consistent with the distinct mechanisms of action for each compound. Utilizing cultured slices from the reporter mice we observed similar selective expression of the GFP in response to the two compounds demonstrating the potential utility of this method for the assessment of proconvulsant properties of new pharmacologic compounds.
1230 Starting Dose Selection for First-in-Human Trials Using Model-Based Approaches.


Selection of the safe starting dose (SSD) in oncology represents a critical decision point in the drug development process, as under- or over-prediction of the SSD can have serious consequences in Phase I, exposing patients to doses that are either subtherapeutic or unsafe. The SSD is commonly estimated by determining if 1/10th the severely toxic dose in rodents (STD10) is tolerated by non-rodents, and if so, using 1/10th of the rodent STD10 dose as the SSD. In general, the STD10 is estimated from GLP toxicology studies where relatively large numbers of animals are divided into a few dose groups to increase statistical power within a group. Here, we propose an alternative method to estimate the STD10 based on logistic regression curve-fitting, (a standard technique for modeling binary outcomes in the clinical setting) with mortality as a binary endpoint. To this end, we simulated mortality data based on 400 different sigmoidal dose-mortality curves with a probabilistic model, and predicted STD10 by logistic regression, comparing this approach to the traditional dose-picking methodology for the same datasets. Model-predicted STD10 values outperformed the traditional method, with better accuracy (absence of downward bias) and precision relative to the true STD10. Additionally, this approach was resistant to the effect of outliers (unexplained mortality). More importantly, the superior accuracy and precision of the STD10 values determined by logistic regression could be achieved with fewer animals used in the conventional approach. A retrospective analysis of an anticancer compound currently in Phase I clinical trials demonstrated potential to narrow the gap between the human Maximum Tolerated Dose and SSD from 110-fold to 10-fold. By using the information from all animals in a study, logistic regression curve-fitting provides the potential for a more robust and less biased SSD, while at the same time decreasing the number of animals utilized.

1231 Application of Electroretinography (ERG) in Early Drug Development for Assessing Ocular Toxicity in Rats.


Retinal ocular toxicity is among the leading causes of drug development attrition in the pharmaceutical industry. Electroretinography (ERG) is a non-invasive functional assay used to assess neuro-ocular physiological integrity by measuring the electrical responses of various retinal cell types. When applied in the pre-clinical setting, ERG may be utilized to evaluate potential ocular toxicity of drug candidates. Studies were designed to assess the sensitivity and specificity of ERG to detect ocular toxicity in Wistar Han rats using several drugs with varied activity in the retina, ranging from no evidence to those with demonstrated microscopic retinal degeneration. To directly assess the utility of ERG, these studies were conducted following a single intravitreal injection (IVT). Doses were selected based on an in-vitro retinal pigment epithelium (RPE) cytotoxicity assay and compound solubility limit. Serial administration of AG-012986 resulted in decreases in b-wave amplitude correlating with increases in plasma levels of retinal toxicity biomarkers (miR-124a and miR-193) as well as marked retinal degeneration as revealed by microscopic evaluation. Here, we propose an alternative method to estimate the STD10 based on logistic regression curve-fitting, (a standard technique for modeling binary outcomes in the clinical setting) with mortality as a binary endpoint. To this end, we simulated mortality data based on 400 different sigmoidal dose-mortality curves with a probabilistic model, and predicted STD10 by logistic regression, comparing this approach to the traditional dose-picking methodology for the same datasets. Model-predicted STD10 values outperformed the traditional method, with better accuracy (absence of downward bias) and precision relative to the true STD10. Additionally, this approach was resistant to the effect of outliers (unexplained mortality). More importantly, the superior accuracy and precision of the STD10 values determined by logistic regression could be achieved with fewer animals used in the conventional approach. A retrospective analysis of an anticancer compound currently in Phase I clinical trials demonstrated potential to narrow the gap between the human Maximum Tolerated Dose and SSD from 110-fold to 10-fold. By using the information from all animals in a study, logistic regression curve-fitting provides the potential for a more robust and less biased SSD, while at the same time decreasing the number of animals utilized.

1232 Absolute Quantitation of Low-Abundance Protein Adducts Using a Novel Accelerator Mass Spectrometry Liquid Sample Interface.


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Drug characterization studies including the detection and quantitation of protein adducts arising from reactive drug metabolites provide important information about a drug candidate’s potential to cause adverse reactions. Current methods for characterizing protein adducts are limited by low sensitivity and difficulty identifying modified proteins. We report the use of a novel accelerator mass spectrometry (AMS) liquid sample interface for absolute quantitation of protein modification by 14C-idoacetamide and identification of peptide modification patterns. Competitive binding experiments showed that covalent adducts of cysteine residues in bovine serum albumin can be quantified at the atomole level in microgram-sized samples of intact protein. Tryptic digests and HPLC separation of labeled peptides followed by peptide analysis using the liquid sample interface and electrospray ionization and tandem mass spectrometry (ESI-MS/MS) showed reproducible patterns of adduct formation, with modifications localized to specific regions of the protein corresponding to known chemical reactivity. Measurements of whole protein adducts using the liquid sample interface were comparable to values measured by standard graphitization with AMS analysis. This technology presents an important novel method for absolute quantitation of 14C-labeled proteins and peptides. Potential applications include micro dosing studies using 14C-labeled protein therapeutics and investigation of post-translational and chemical modifications of proteins.

1233 Evaluation of Respiratory Function in the Conscious, Nonrestrained Cynomolgus Monkey Using Respiratory Inductive Plethysmography.


Evaluation of the potential alteration of respiratory function is mostly performed in rodents using whole body plethysmography, while cardiovascular telemetry studies are usually conducted in large animals. The purpose of the present study was to evaluate the use of respiratory inductive plethysmography (RIP) as a method that might be suitable for routine evaluation of respiratory function in primates. Four monkeys were equipped with jacketed external telemetry devices including abdominal and thoracic belt sets (JET, DSI). Thoracic and abdominal signals were recorded and analysed using Premam closed software. After an acclimation period, animals received theophylline (100 mg/kg, p.o), clonidine (100 μg/kg, i.m) and corresponding vehicle (p.o or NaCl 0.9%, i.m). Respiratory rate (RR), tidal volume (TV) and minute volume (MV) were recorded continuously over 24 hours. Theophylline induced increases in TV, MV and RR when compared to vehicle (Emax: +59%, +118% (p < 0.01) and + 41% (p < 0.01), respectively). Clonidine produced significant decreases in TV and MV and RR (Emax: -12%, -29 % (p < 0.01) and -27% (p<0.05)). The results demonstrate the validity of the RIP method for the assessment of respiratory function in the monkey for a long period (24 hours). These findings support the use of the RIP method for combined assessment of potential effects of drug candidates on cardiovascular and respiratory functions. The methodology could allow the integration of respiratory investigations into regulatory toxicology studies in primates. Such an approach is particularly applicable for the safety evaluation of biotechnology-derived products, for which rodents are often not applicable models.

1234 Impact of Food and Fecal Contamination on Dog Urinalysis Data.


Suspected aberrant urinalysis results were observed in dogs urinating limited urine volumes, or presenting clinical diarrhea, prompting an investigation to determine potential causes of contamination. Diet consumed and feces were identified as potential sources. Dietary effects on blood urine contents were determined using two laboratory diets: Certified Canine Chow No. 5007 and Harlan T eklan Certified 25% Lab Dog Diet (2025). Urine was collected from animals deprived of food and water, previously fed either diet during a 5 hour period as well as following an overnight collection period. Each diet was mixed with saline to mimic contamination. Five hour small urine samples collected from animals fed diet 5007, and 5007 diet-saline suspensions, both tested positive for moderate (3+) to large (4+) amounts of blood measured semi-quantitatively with urine dipsticks (Multistix 14SG, Siemens). 2025 diet-saline suspensions tested positive for traces of blood. Larger volumes and more dilute urine collected following overnight collection tested negative for blood indicating levels were below the detection threshold. The effect of fecal contamination, in dogs presenting diarrhea, on several markers of kidney function was also evaluated. Varying amounts of feces, from 1 to 5 grams were added to approximately 8 mL urine to mimic contamination. Contaminated and uncontaminated samples were analyzed for N-Acetyl-B-D-glucosaminidase (NAG), Gamma-glutamyl-transferase (GGT) and protein, with values corrected for urinary creatinine concentration. Increases in kidney markers were proportional to the amount of feces added.
Simultaneous modulation of epidermal growth factor receptor (EGFR) and insulin like growth factor receptor 1 (IGF-1R) signaling, as an oncology therapeutic strategy, is based on cross talk between the two signaling pathways to overcome resistance developed if only a single pathway is targeted. However, evidence of class-related adverse effects of individual anti-EGFR and anti-IGF-1R agents in humans including diarrhea, acne form skin rash and interstitial pneumonitis (EGFR-specific) and hyperglycemia, thrombocytopenia and Cardiovascular toxicities (IGF-1R-specific) has emerged in clinical and non-clinical toxicity studies. In contrast to marketed anti-EGFR monoclonal antibodies which bind to domain III of the extracellular portion of EGFR, BMS 964210 binds with high affinity to domain I and overlaps the EGF and TGFβ binding pocket of both human and cynomolgus monkey EGFR and IGF 1R. In a 4 week intravenous (2QW) toxicity study with BMS-964210 in cynomolgus monkeys (6, 12 and 25 mg/kg), pharmacodynamic elevation of plasma biomarkers of EGFR and IGF 1R blockade (TGFα, amphiregulin, and IGF 1) were observed. Despite the generation of anti BMS 964210 antibodies, systemic exposures following first dose were dose proportional and elimination was linear. No evidence of rash, pulmonary findings, hyperglycemia or thrombocytopenia was noted, and there were no drug related deaths or effects on neurologic, respiratory, or ophthalmologic endpoints. At the highest dose, two female monkeys suffered body weight loss and dehydration that required fluid administration for 3 days. Based on the highest non-severely toxic dose (HNSTD) of 12 mg/kg, there is sufficient safety margin to the proposed clinical starting dose of 0.5 mg/kg (QW). These preliminary data suggests that BMS-964210 may have an improved tolerability profile compared to marketed anti-EGFR mAb and deserves further evaluation in clinical studies.

1235 High-Resolution Isotope Dilution (HRID) Quantitative Analysis: Metabolites in Safety Testing (MIST) Application to Diclofenac.

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During early clinical development it is important to understand the identity and amount of circulating metabolites in man. Are there human specific metabolites? Guidelines have been issued by regulatory agencies to address this question (CDER, 2008). We are interested in applying the metabolism of isotope labeled drugs [14C alone or both 14C and 13C] and HRID to accurately determine MIST liability in the clinic in a manner that is both rapid and not excessively expensive (e.g., involve AMS, de novo synthesis and bioanalytical method development for relevant drug metabolites). Simplicity and ruggedness are also important aspects of these efforts. This work investigated diclofenac (D), Diclofenac, [14C]D (62.7 mCi/mmol) and [13C6]D were purchased. Human hepatocytes were purchased from Life Technologies, Grand Island, NY. Incubations were conducted following Life Technologies procedures. Chromatographic separation was achieved using an Agilent 1200 HPLC system and detection of radioactivity was by an IN/US β-Ram Model 3. MS analysis was by Thermo LTQ Orbitrap Discovery MS (+ ESI). No special tuning was done for D metabolism analysis. Since the specific activity of the label was known and fixed, the amount of drug metabolite/parent drug in the incubates at 60 minutes were determined using LC-RAM. Assuming that the amounts of drug metabolites produced from [13C6]D was similar to the amount produced for the [14C6]D allowed for the construction of standard curves for all drug-related material within the limits of detection. This allowed for the quantitative analysis of drug-related material including, 4-hydroxy-D and D-glucuronide. The method resulted in acceptable linearity, sensitivity and reproducibility for all analytes. A second key concept is that using HRMS compared to MS-MS using tandem quadrupole affords considerable advantages in simplicity and ruggedness, a key objective.

1236 12-Week Intrathecal Administration Study in Port Catheterized Juvenile Cynomolgus Monkeys.

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The cynomolgus monkey is the predominant NHP species when it comes to preclinical safety evaluation of new medical products. Assessment of jv. toxicity within this species represents an emerging field, requiring established techniques for 12 month or younger monkeys. The objective of the study was to determine the feasibility of bi-weekly bolus delivery in an intrathecally implanted port catheter system to the jv. monkey for 12 wks and to assess the feasibility of CSF collection. Vehicle or PBS was administered to 6 implanted (between L2-L5) monkeys (8-13.5 months; ~800 g). Clinical signs, bw, neurological examinations, CSF, and clinical pathology data were obtained. Perfusion necropsy, spinal cord brain trimming and pathology were conducted. There were no indications for procedural-related clinical signs, bw development, clinical pathology or neurological findings. Increased AST was observed on day 2, most likely related to manual restraint of the animals on day 1. Necropsy confirmed the placement of the catheter tips at T11/12 in 4/6 animals. There were no adverse effects noted at the site of the catheter in the intrathecal space and in particular, no evidence of pronounced reactions at the catheter tips. At the catheter tip, tissue reactions were consistent with those reported to occur with the placement of intrathecal catheters in multiple species (M Butt, 2011). Changes at the catheter tip included slight to minimal fibrosis, adhesion to the overlying dura, and slight compression of the spinal cord (no cord damage). Overall lumbar CSF sampling through the port was only possible in 1/6 animals. However, spinal lumbar CSF sampling under sedation using a Pennan Paed® needle proved to be successful on all occasions. Obtained CSF indicated a low level of white blood cells after surgery, and minimum of red blood cells contamination. In conclusion, lumbar intrathecal administration of juvenile cynomolgus monkeys for up to 12 wks using a surgically implanted port catheter system is considered feasible.

1237 BMS-964210, a Pegylated Bispecific Adnectintargeting EGFR and IGF-1R, Demonstrates Improved Class-Specific Toxicity Profile in Cynomolgus Monkeys.

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At our Laboratory, the microscopic examination of subcutaneous injection sites involves the evaluation of three longitudinal sections per tissue sample. We conducted a retrospective study to evaluate the relevance of the diagnoses in subcutaneous injection sites if only one section is examined. Two rat studies and three primate studies were reviewed. All animals received saline or test item injected subcutaneously in the dorsum. At necropsy, 3 pieces, each 3 mm wide, were trimmed from each site. Piece 1 was in the middle, piece 2 lateral to it and piece 3 medial to it. The original study pathologists formulated overall diagnoses for each site, taken from evaluation of each of the 3 pieces. One pathologist reviewed the three sections from each site with reference to the findings recorded by the original study pathologists and attributed each of the original diagnoses to the section(s) where it was observed. In each species, animals receiving saline or test item were evaluated separately. Statistical analysis was performed, comparing section 1 with the other two sections and with the original overall diagnosis. In the rat, section 1 had a statistically significantly higher number of findings when compared to other sections, in one species and in one animal. In the primate, section 1 did not differ from the other sections in control and treated animals. In both species, however, section 1 had a statistically significantly lower number of findings when compared with the original overall diagnosis. Contrary to the results we obtained in a previous study concerning intramuscular sites, this retrospective study reveals that examination of three sections of subcutaneous injection sites improves the quality of diagnosis. For microscopic changes induced by subcutaneous injections, particularly in primates, one middle piece is not considered to be sufficiently representative of the whole site.
Internal Exposure of Common Environmental Pollutants in a Chinese Population.

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Studies indicated that exposure to environmental pollutants could increase the potential risk of human’s health. Many efforts have been made to detect the environmental pollutants, but such data cannot be of concern to ordinary people for no one knows how many environmental pollutants and at what amounts were exposed to themselves. In order to provoke the attention of public and point out the individual who is at risk mostly, it is necessary to measure the internal exposure of environmental pollutants. These data can directly reflect the exposure of those pollutants to ordinary people. In present study, we measured the total urinary concentrations of eleven common pollutants in a Chinese population by GC, GC-MS, HPLC and ELISA. We found the percentage of people with detectable levels of an individual chemical ranged from 13% to 100%. Cotinine (COT) was detected in 88% of the persons (n=2100), dibutyl phthalate (DBP) was detected in 97% of the persons (n=800), the detection rate of aflatoxin (AFT) and toluene (TOL) were 85% and 13%, respectively (n=600); organophosphorus (Org-p) was detected in 99% (n=400), tartrazine (TAR) was detected in 78% (n=351), the detection rate of diethylstilbestrol (DES), methanolic acid (MA), Styrene (STY), terachlorodibenzo-p-dioxin (TCDD) and nonyl phenol (NP) were 100%, 100%, 96%, 32% and 21%, respectively (n=100). The 95th percentiles of COT, DBP, AFT, TOL, Org-p, TAR, DES, MA, STY, TCDD and NP were 44 ng mL-1, 9 ng mL-1, 39 ng mL-1, 39 ng mL-1, 0.28 ng mL-1, 33.92 ng mL-1, 22, 000 ng mL-1, 23, 000 ng mL-1, 930 ng mL-1, 1,600 ng mL-1, 0.33 ng mL-1, respectively. The present data show that people in China are widespread exposure to multiple pollutants. Therefore, in order to describe the real risk of individual, a well designed study is needed to evaluate the associations between internal exposure of those pollutants and potential adverse health consequences.

Occurrence of Gaseous Pesticides and Other Hazardous Volatiles in Sea Containers Arriving in Sweden.

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Individuals who enter containers for inspection, unloading or cleaning may be exposed to gaseous pesticides (fumigants) and other volatiles. Previous reports from Hamburg and Rotterdam, as well as a few incidents in Sweden, have raised concerns about the exposure situation in Swedish container terminals. Since systematic studies in Sweden are missing, we performed a pilot study to investigate the occurrence of hazardous volatile chemicals in import containers. Air was sampled from 101 randomly selected containers in the port of Gothenburg. Air samples were drawn without opening the containers, as they passed and briefly halted at the import inspection station, and analysed by FTIR spectrometry. One container (1%) contained detectable residues of fumigant (1 ppm carbaryl sulphide), although none was labeled as fumigant treated. As in the two previous studies, a number of other volatile chemicals were detected. The most common ones were methanol (78%) and carbon monoxide (45%), with all readings below the occupational short term excursion limits (STEL). These substances are likely degradation products from the plywood flooring as they were also detected in empty containers. Other frequently detected chemicals include hydrocarbons (unspecified, 47%) and ammonia (15%), with some readings above or well above the STELs or ceiling limits (CL). In our judgement, based on the present and two previous studies, the probability of life-threatening exposure is low. However, violations of STELs and CLs do occur, thus prolonged or repeat visits in unchecked containers may constitute a significant health hazard. In conclusion, extensive and systematic investigations of import and export containers are greatly needed as basis for risk management. Methods for rapid and efficient ventilation should be developed. Prior to entering it should be ensured, by adequate measurements and/or ventilation, that the container is safe.

Interactions between Urinary 4-Tert-Octylphenol Levels and Metabolism Enzyme Gene Variants on Idiopathic Male Infertility.

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Background: Octylphenol (OP) and Trichlorophenol (TCP) are the representative members of alkylphenols and chlorophenols, which act as endocrine disruptors and have effects on male reproductive function. Objectives: We studied the interactions between 4-tert-Octylphenol (4-t-OP), 4-n-Octylphenol (4-n-OP), 2,3,4-Trichlorophenol (2,3,4-TCP), 2,4,5-Trichlorophenol (2,4,5- TCP) urinary exposure levels and polymorphisms in selected xenobiotic metabolism enzyme genes among 589 idiopathic infertile male patients and 396 controls in a Han-Chinese population.

Methods: Ultra high performance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS) was used to measure alkylphenols and chlorophenols in urine. Polymorphisms were genotyped using the SNPstream platform and the Tagman method. We used likelihood ratio tests (LRT) to explore these gene-environment interactions in idiopathic male infertility, and used false discovery rate (FDR) to adjust for multiple testing.

Results: Among four phenols that were detected, we found that only exposure to 4-t-OP increased the risk of male infertility (P trend=1.70×10-7). The strongest interaction was between 4-t-OP and rs918758 in CYP2C9 (P trend=6.05×10-7). It presented a significant monotonic increase in risk estimates for male infertility with increasing 4-t-OP exposure levels among men with TUGCC genotype (low level compared with non-exposed, odds ratio (OR) =2.26, 95% confidence intervals (CI) =1.06, 4.83; high level compared with non-exposed, OR=9.22, 95% CI=2.78, 30.59; but no associations observed among men with TT genotype). We also found interactions between 4-t-OP and rs986894 in CYP2C19, and between rs1038943 in CYP1A1, on male infertile risk (P trend=8.05×10-7, 3.73×10-4, respectively).

Conclusions: We observed notable interactions between 4-t-OP exposure and metabolism enzyme gene polymorphisms on idiopathic infertility in Han-Chinese men.

Smartphone Technologies Coupled with Environmental Models Allow for Large Scale Community Outreach and Risk Assessment.

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Chronic exposure to several air pollutants has been identified by the EPA as a major source of concern for human health. Environmental models have been created to predict spatial distributions of air pollutants, but are unable to account for daily travel among the general US population. Several GPS-based methods of tracking a person’s movement through areas of concern have been developed, but require expensive or bulky GPS units, have limited ability to provide timely results to participants, and the accuracy of GPS is not well maintained across all demographic groups, with more than 100 million US smartphone users. We developed in-house and android applications with GPS functionality to predict personal and population-wide exposures to fine particulate matter (PM2.5), coarse particulate matter (PM10), and ozone. Environmental distribution models were created using Kriging algorithms with MODIS satellite imagery and Oregon Department of Environmental Quality hourly PM2.5, PM10, humidity, ozone, and temperature measurements. Geographic locations were sampled from four smartphone devices at 30 minute intervals (n=10000 sampling events). Locations and corresponding times were sent to a database running the environmental modeling software. Predicted exposure levels, times, and locations were returned to corresponding smartphones and were anonymously added to a data set of predicted exposures collected from all participants. Personal exposure levels were presented to participants as interactive smartphone maps and graphs. Hotspot exposure sites were predicted by partitioning Oregon into 10 km2 regions and mapping daily group exposure levels within each region. This project was created with open-source software and can be scaled to larger groups with minimal cost. This research was supported by NIH P42 ES014465.
Toxicology studies are moving to high-throughput methodologies, however, often times the chemical concentrations are not analyzed or verified. Historically, analytical methods capable of measuring organics in aqueous solutions (including fish water and cell media) often relied on more complicated techniques. The technique presented here has reduced 99% of the historically used sample preparation. Thereby reducing the time per sample associated with preparation from ~90 minutes to <1 minute. This technique essentially eliminates the chance for target analyte loss during sample preparation and therefore eliminates the need for surrogate standards, such as 13C-labeled, which is typically used to correct for analyte loss during sample preparation. This method utilizes an innovative liquid-liquid extraction, where ~10-25 μL of cell media or water is transferred to a gas chromatography (GC) vial with 1 mL of hexane and 50 mg of sodium sulfate. Target analytes partition from the water to hexane and the water is captured by the sodium sulfate. Overall extraction efficiency is ~100%. The utility of the novel method was demonstrated with a proof of concept experiment that evaluated BDE 47 water concentrations overtime from glass and plastic (48-well plate) wells. BDE 47 concentrations measured in the glass well remained consistent over 24 hrs; however there was a rapid decrease in BDE 47 water concentrations (~80%) within the first 8 hrs in the plastic well. These results demonstrate that this method is simple and useful for both water and cell media studies and offers the ability to monitor the same well over time. Many exposures in toxicology studies start at concentrations above solubility and are used in plastic plates. While this has been an acceptable method of rapidly obtaining LC and IC50 values, it may not be an accurate representation of the actual water concentration. By accurately analyzing the concentration and determining the uptake, more accurate results may be obtained and used for risk assessment and mechanistic determinations, while adding increased study to sample comparability.

1244 Evaluating High-Throughput Exposure Models with NHANES Data.


Prioritizing screening and testing for large sets of chemicals for potential risk requires rapid exposure assessment, ideally based on chemical properties that are easily measured and/or computed. There are currently few such approaches, but like any predictive methodology these need to be evaluated against real exposure data. We evaluated two models, “Risk Assessment, Identification and Ranking” (RAIDAR; www.armresearch.org/index.html#!/page_RAIDAR_DL) and USEtox™(www.usetox.org) against the National Health and Nutrition Examination Study (NHANES) 2005 and 2011 data on urine concentrations of xenobiotics. Using NHANES to evaluate exposure models requires addressing several issues: chemicals measured in urine need to be linked back to potential exposures, and there may be multiple relationships between parent chemicals to which people are exposed and the product chemicals measured in urine; many chemicals were below limits of quantitation for a large fraction of the sampled subjects; variation among chemicals is a function of variation of exposure, pharmacokinetic (PK) variability, and the distribution of relative times of exposure and sampling. Steady state PK assumptions and a Bayesian statistical framework were used to infer parent exposures from NHANES urine products. We further use Bayesian statistical methods to evaluate the reduction of median population exposure to the parent chemicals on RAIDAR and USEtox predictions, estimated production volume, and simple indicators of household use. Household use was the single most important factor. Among those chemicals with a consumer use, median exposure was correlated with production volume, but there was little evidence of a relationship between estimated exposure and any of the predictors among chemicals with no household use. The resulting model can be used for predicting exposure potential for new chemicals, along with the uncertainty of that prediction. Refining what constitutes ‘household exposure’, as well as including more exposure datasets, should reduce the uncertainty of the current model. [This abstract does not necessarily reflect EPA policy.]

1245 Haematology and Erythrocyte Osmotic Fragility Indices in Domestic Chicken following Exposure to a Polyvalent Iodophorous Disinfectant in Water Treatment.

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The effect of prolonged use of lodosteryl, a polyvalent iodophorous compound, as water disinfectant, on the haematology and erythrocyte osmotic fragility of the domestic chicken was investigated. Twenty-eight adult male domestic chickens of the Nera black strain were divided into four groups of seven birds per group. Birds in groups B-D were given potable water containing 1 ml, 2 ml and 4 ml/l lodosteryl respectively for six weeks. Group A served as the control. Blood samples were collected from each bird after six weeks and analyzed immediately. No significant changes were observed in the packed cell volume (PCV), haemoglobin (HB), mean corpuscular volume (MCV), mean corpuscular haemoglobin (MCH), mean corpuscular haemoglobin concentration (MCHC), platelet total and differential leucocytes values. However, red blood cells (RBC) were slightly lower while erythrocyte osmotic fragility and erythrocyte sedimentation rate (ESR) was higher in those birds exposed to lodosteryl compared with control. This study confirms that prolonged use of lodosteryl is stressful and may lead to intravascular haemolysis as indicated by the higher erythrocyte fragility and ESR values, respectively. The damage observed, may be due to peroxidation of erythrocyte membrane lipids, proteins by generation of free radicals induced by iodine.

1246 How Inaccurate are Serum-Lipid Unadjusted Organochlorine Levels?


Introduction. Organochlorine pesticides (OCs) have been banned by many countries, but they are still detected in human and animal tissues worldwide. An empric assessment comparing serum lipid adjusted and unadjusted OC levels has been, so far, not reported in the literature. Objective. To estimate how inaccurate are lipid-unadjusted OCs serum levels compared to adjusted ones. Methods. In a survey carried out in a rural population in Brazil exposed to OCs since early 1960’s, blood samples were obtained from 995 residents of both sexes and all ages. Serum concentrations of 19 OCs were determined by gas chromatography with electron capture detection and were further adjusted by triglyceride and cholesterol content. Pearson correlation coefficients and determination coefficients (R2) of serum lipid-adjusted and unadjusted concentrations of p,p’-DDE, beta-HCH and aldrin were calculated. Results. Among male and female adults (> 14 yr.), median (1st and 3rd quartiles) of serum-lipid adjusted concentrations were, respectively: 8.32 (2.86-21.9) and 9.64(3.45-28.9) for p,p’-DDE; 6.0 (2.08-15.4) and 6.98 (2.81-17.6) for beta-HCH; 1.89 (0.73-11.0) and 2.44(0.77-14.1) for aldrin. Determination coefficients of serum lipid-adjusted and unadjusted OCs were R2= 0.91 for p,p’-DDE, R2= 0.90 for beta-HCH, and R2= 0.93 for aldrin. Correlation between serum lipid-adjusted and unadjusted concentrations was indeed higher for those OCs with serum levels < 50 ng/ml. Conclusion. When serum lipid content is unavailable, unadjusted OC levels can be reasonably used to estimate internal dose of these chemicals, particularly when concentrations are < 50 ng/ml.


Surface deposition of insecticides applied as indoor residential foggers, baseboard or perimeter sprays, spot sprays and crack-and-crevice sprays represent pathways of unintentional, and unavoidable post-application exposure of children and adults. Estimation of the magnitude of this exposure following an application event is associated with uncertainty due to many factors including 1) surface residue deposition and distribution, 2) access to and the nature of contact with treated surfaces based on time-activity patterns of residents, and 3) the role of residue removal mechanisms such as cleaning treated surfaces, pesticide degradation or redistribution, hand washing and bathing following contact. A comparative spatial deposition study was conducted involving indoor residential fogging, perimeter and crack and crevice application methods. Residues measured using a spatial grid of deposition dosimeters on floor surfaces demonstrated significantly lower residue concentrations in readily
Our data suggest that disturbed placental angiogenesis, via upregulation of sFLT1, may link prenatal arsenic exposure to increased risk on SGA.

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**1250** Plasma Polybrominated Diphenyl Ethers (PBDEs) in Californian Women at High Risk for Birthing an Autistic Child.

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Exposure to the polybrominated diphenyl ethers flame retardants (PBDEs) is a major preventable health concern. Little is known about the extent and patterns of PBDE exposures during pregnancy, especially in populations susceptible to heritable neurodevelopmental disorders. We measured plasma PBDE levels in Californian women participating MARBLES (Markers of Autism Risk in Babies–Learning Early Signs) who are at high risk for birthing an autistic child. BDE-28, -47, -49, -52, -95, -99, -100, -136, -153, and -183 were measured using GC/MS/MS in plasma samples collected from 79 women at each trimester and at delivery (251 samples total). PBDEs were normalized to plasma volume (ng/ml) and total lipids (ng/g). The concentrations of maternal PBDEs in MARBLES were compared to data from the National Health and Nutrition Examination Survey (NHANES 2003-2004). All ten congeners were detectable in the maternal samples from MARBLES, with BDE-100 (0.792 ± 0.540 ng/ml, 163.5 ± 117.5 ng/g) and BDE-47 (0.061 ± 0.458 ng/ml, 141.1 ± 117.6 ng/g) contributing the highest abundance. Compared to NHANES, women enrolled in MARBLES had significantly higher mean concentrations of BDE-47 (23.9 versus 141.1 ng/g), -99 (5.51 versus 92.3 ng/g), -100 (6.06 versus 163.5 ng/g) and -153 (3.90 versus 28.7 ng/g). Plasma PBDE (ng/ml) in women 25 to 35 years of age increased during gestation, but decreased if they were older than 35 years. Because of the increase of total plasma lipids during gestation, PBDE expressed on an ng/g lipid basis decreased in all age groups with gestational stage. Fluctuations in maternal plasma PBDE levels during pregnancy illustrate the importance of gestational age and underscore the non-linear relationship between volume- and lipid-corrected PBDE concentrations.

**1251** Estimation of Dietary Lead Exposure for US Children Using a New Method: Implications for the Integrated Exposure Uptake Biokinetic Model for Lead in Children.

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In addition to site-specific inputs, the Integrated Exposure Uptake Biokinetic Model for Lead in Children (IEUBK model) uses national defaults that are recommended when site-specific information is not available. As part of a periodic evaluation of inputs to the IEUBK model, the methodology underlying the basis for default calculations of dietary lead intake has been updated. Previously, the food consumption values in the IEUBK model were based on information reported by Pennington (1983) and the lead in food values were based on food residue data from the ongoing FDA Total Diet Study; the latter are updated periodically as new data become available from FDA.

Recent advances in methods for estimating dietary intake provide a current and scientifically sound basis to develop nationally-representative, age-specific values for food consumption. The dietary component of NHANES, called the What We Eat in America (WWEIA) Dietary Survey, includes two 24-hour dietary recall interviews during which each respondent reports the amount of all foods they consumed during the prior day. The food consumption values (grams/day) were estimated using a non-linear mixed model developed by the National Cancer Institute (the NCI method). The NCI method uses information on meal sizes and frequency (probability) with which specific food items are consumed to estimate daily consumption rates. The NCI method produced estimates of dietary lead intake that were 18% higher than the previously estimated values, depending on the food item. Further analysis revealed the increase in the dietary lead intake values was largely due to an increase in the estimated daily consumption values rather than higher lead concentrations in food.
The default values in the IEUBK model (Version 1.1, Build 11) are based on values reported in the Office of Air Quality Planning and Standards report (U.S. EPA, 1989) and the IEUBK Model Technical Support Document (U.S. EPA, 1994). More recent data provide a more scientifically sound basis to further develop nationally-representative, age-group specific values for ventilation rates in children. The default values in the IEUBK model were derived based on body size in combination with smoothed data from Phalen et al. (1985). EPA's (2008) Child-Specific Exposure Factors Handbook provides recommendations for long-term (>30 days) ventilation rates that are based on the average of several studies (Arcus-Arth and Blaisdell, 2007; Brochu et al., 2006; Stiefelman, 2007). Because Arcus-Arth and Blaisdell (2007) used indirect measures of ventilation rates based on dietary and activity survey responses, it was not considered acceptable for derivation of default values for the IEUBK model. Brochu et al. (2006) and Stiefelman (2007), however, were based on the doubly-labeled water (DLS) energy data from the Institute of Medicine. DLS energy data are recognized as the gold standard for energy expenditure and an improvement over ventilation estimates based on dietary recall or activity survey data.

Ventilation rate was calculated from total energy expenditure using Layton's approach as described by Stiefelman (2007). The analysis was on pooled data for males and females. More detailed analysis of ventilation rates as a function of age and gender showed the estimated ventilation rates to be parallel and 7% greater in males than females. The resulting ventilation rate values are between 19-66% higher than the existing IEUBK model defaults. Because these values are derived from energy expenditure information, they provide a more scientifically defensible basis for the default values in the IEUBK model.

Exposure Conversion Factor and Discretized Bayesian methods. The means of the estimated dose distributions were comparable to average daily doses estimated using other methods (point estimates), such as food concentrations multiplied by a consumption rate. Probabilistic reverse dosimetry, however, also provides the distribution of smoothed data from Phalen et al. (1985). EPA's (2008) Child-Specific Exposure Factors Handbook provides recommendations for long-term (>30 days) ventilation rates that are based on the average of several studies (Arcus-Arth and Blaisdell, 2007; Brochu et al., 2006; Stiefelman, 2007).

First, Monte Carlo simulations of a physiologically based pharmacokinetic (PBPK) model were performed to account for variability/uncertainty in exposure factors and PK. Next, the simulated exposure-biomarker relationship was used to convert urinary perchlorate concentrations to a distribution of daily doses. The conversion was conducted using a web-based tool, Probabilistic Reverse Dosimetry Estimating Exposure Distribution (PROceed), with two methods: the Exposure Conversion Factor and Discretized Bayesian methods. The means of the estimated dose distributions were comparable to average daily doses estimated using other methods (point estimates), such as food concentrations multiplied by a consumption rate. Probabilistic reverse dosimetry, however, also provides the distribution of dose estimates to represent the variability/uncertainty in exposure factors and PK. [This abstract has been cleared by the EPA but solely expresses the view of the authors]
1257 Lipidomics of Subchronic Low Level Cadmium Exposure in the Rat.


Epidemiological studies suggest an association between cadmium in drinking water and vascular diseases. However, the precise cadmium mechanism of action remains enigmatic. This study was undertaken to investigate the effect of cadmium on lipid metabolism of Wistar male albino rats by exposing the animals to 100, 200 and 300 ppm cadmium doses for 12 weeks in their drinking water. Control animals received distilled water for the same period. At the end of 12 weeks, dyslipidemia induced by the cadmium doses exhibited different patterns. Dose-dependent hypolipidemia and hypotriacylglyceridemia characterised the effect of cadmium exposure at all doses whereas plasma free fatty acid (29%) was increased by cadmium exposure. Reverse cholesterol transport was inhibited by all the cadmium doses as evidenced by 65% decreased HDL cholesterol concentrations whereas hepatic cholesterol was decreased by 55%. Renal and brain cholesterol (46%, 65%) and triacylglyceride (62%, 50%) were dose-dependently decreased by cadmium exposure respectively; on the other hand, exposure to cadmium decreased cardiac cholesterol by 45%, but enhanced/balanced triacylglyceride content. Cadmium at all doses of exposure inhibited both hepatic and brain HMG CoA reductase by 49% and 61% respectively. We observed positive association between tissue cadmium levels and plasma FFA, and negative associations between tissue cadmium levels and HDL cholesterol. Our findings indicate that in contrast to strengthening a dose-dependent effect phenomenon as observed with many other compounds, cadmium up- or down-regulate different pathways in the lipid metabolism spectrum at “low” or “high” doses and this might be responsible for the insidious vascular effects.

1258 Serum Great Lakes Pollutant Levels in Lake Ontario Fish and Wildlife Consumers.

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Industrialization of the Great Lakes region resulted in the discharge of many halogenated aromatic hydrocarbons (HAHs) into the Lake Ontario basin. Due to their persistence and lipophilicity, HAHs can biomagnify in the food chain and are a potential source of human exposure to polychlorinated biphenyls (PCBs), polychlorinated dibenzo-p-dioxins (PCDDs), dibenzofurans (PCDFs) and pesticides. Consumption of Lake Ontario fish and wildlife is one factor that contributes to higher serum levels of persistent HAHs and potentially greater risk of adverse human health effects. The New York State Angler Cohort Study (NYSACS) is a large prospective cohort of licensed anglers from 16 New York counties in close geographic area (n=16). We examined serum levels of 20 PCBs, including PCB 126, 153, 138, 180 and 170. The consumer group had significant elevated levels (n=27) and with self reported consumption of Lake Ontario fish were chosen, and matched to subjects having never consumed Lake Ontario fish by age, sex and pesticide levels. For this analysis, participants with the highest serum PCB153 levels (n=27) and with self reported consumption of Lake Ontario fish were chosen, and matched to subjects having never consumed Lake Ontario fish by age, sex and geographic area (n=16). We examined serum levels of PCBs, PCDDs, PCDFs and pesticides by GC/ECD and GC/MS. The consumer group had significant elevated levels of 20 PCBs, including PCB 126, 153, 138, 180 and 170. The consumer group also had a 1.6-fold increase in 2,3,7,8-TCDD, a 1.4-fold increase in 1,2,3,7,8-PentaCDD, a 1.54-fold increase in 2,3,4,7,8-PentaCDF, a nine-fold increase in mirex, a 2.5-fold increase in DDE and a two-fold increase in t-nonachlor relative to the non-consumer group (all values p<0.05). Together, the results indicate that consumption of fish and wildlife from Lake Ontario is a source of exposure to mirex and other HAHs. (Supported in part by ATSDR, Grant H75-ATH 298338)

1259 Exposure to Environmental Tobacco Smoke Causes Endotoxin Tolerance.

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Exposure to cigarette smoke is known to increase susceptibility to and severity of pulmonary diseases such as bronchitis and emphysema. During tobacco manufacturing processes, in which the temperature and humidity are brought to an optimum level for fermentation, tobacco is colonized by fungi and bacteria. Lipopolysaccharide (LPS), a gram negative bacterial component, has been found in tobacco smoke. In this study, we wanted to assess whether acute exposure to environmental tobacco smoke (ETS) could alter the immune response caused by exposure to the bacterial endotoxin, LPS. Using a C57BL/6 mouse model, we compared the inflammatory cytokine levels secreted by LPS stimulated alveolar macrophages in ETS exposed and unexposed (control) groups. Tumor necrosis factor-alpha (TNF-a) levels were significantly attenuated in the ETS exposed groups after ex-vivo exposure to LPS in comparison to the control group. We observed that whether the ex-vivo LPS exposure was performed immediately after the ETS exposure or 24 h after the ETS exposure, LPS-induced TNF-a expression was significantly reduced by ETS exposure. No significant difference was observed in alveolar macrophage recovery between groups. Also, no significant difference was found in cell viability between the two groups. This suggests that TNF-a attenuation in ETS exposed groups resulted from endotoxin tolerance caused by LPS or LPS-like constituents in ETS. This endotoxin tolerance resulting from acute ETS exposure can suppress the immune function and subsequently increase susceptibility to bacterial infections. These results provide insights into the weaker immune defense observed in smokers and secondhand smokers. This study was supported by the Flight Attendants Medical Research Institute grant 072083.

1260 Mitochondrial-K+-ATP Channel Activation Rescues Phosgene-Induced Toxic Inhalational Lung Injury.

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To determine whether R-801, a mitochondrial-K+-ATP channel activator, is effective in rescuing acute lung injury (ALI) induced by phosgene (COC12), C57BL/6 mice were exposed to COCl2 (5 ppm) or filtered air (FA) for 20 min. Animals were administered intraperitoneal (IP) R-801 (80 mg/kg/dose) or vehicle (hydroxypropyl cyclodextrin, HPDC) at 2 and 6 h after COCl2. Animals were tracheostomized, bronchoalveolar lavage (BAL) performed, and lungs harvested at 24 and 48 h. COCl2-exposed animals developed severe lung injury compared to controls, with increased interstitial and alveolar edema, hemorrhage, and recruitment of inflammatory cells. Treatment with R-801 led to significant reduction in histologic lung injury and near normalization of lung histology Immunostaining for Ly-6G (Gr-1) at 48 h demonstrated significant neutrophil (PMN) accumulation around bronchovascular bundles in COCl2-exposed animals which was profoundly attenuated in R-801-treated COCl2-exposed animals. Furthermore, levels of the pro-inflammatory cytokines and BALF protein were dramatically reduced compared with controls. Treatment with R-801 restored epithelial barrier integrity after COCl2 with striking attenuation in BALF protein levels 48 h after COCl2. Taken together, our findings demonstrate that selective mito-K+-ATP channel opening with R-801 is highly effective at rescuing COCl2-induced ALI via attenuation of inflammation, restoration of epithelial tight junction barrier integrity, and induction of endogenous surfactant protein expression.

1261 Systematic Review of BPA “Low Dose” Literature in the Context of Human Dosimetry Exposes a Need to Set Standards for Responsible Communication of Both Toxicity and Exposure Data.

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BPA is a weakly estrogenic monomer used for making polycarbonate plastics and food packaging liners. At sufficient exposures, BPA is toxic in rodent non-human test systems. Uncritical reference to many toxicity studies as “low dose” has led to the belief that exposure levels in these studies are similar to human exposure levels.
implying that BPA is toxic to humans at current exposures. However, a comprehen-
sive assessment of exposures in humans and test systems has not been con-
ducted. Applying the fundamental principles of biomonitoring, exposure assess-
ment and dosimetry, we conducted a systematic review of BPA exposure levels in
130 peer-reviewed in vivo and in vitro BPA toxicity studies self-referring as “low
T.
2.5 ppm amine or the equivalent concentration of degraded amines
3. Inflammatory response in lungs was assessed by counting inflammatory cells in
4. Laboratory-generated degradation products formed via addition of CO2 and O2 were
5. Carbon dioxide (CO2) adsorption with aqueous amine solvents is among the lead-
6. This study evaluated the inflammatory response associated with inhalation expo-
7. The SPHERES research program was created to examine the role of inhalated sec-
8. The only SOA exposure in which a statistical increase in TBARS was measured,
9. The acidic α-pinene mixture, as well as α-pinene + NH3 (both neutral and acidic) and
10. Alkaline Comet assay whereas the increase of oxidized base (8-OHdG) was 1.1 fold
11. In workers who were smokers, there was a continuously increased over the years. Manufactured goods made of polymers are generally complex materials. These are composed of polymers or copolymers them-
12. The importance of plastic materials for different applications in everyday life has con-
13. The relationship between these parameters in petrochemical plant workers during a
14. The SPHERES Program: Secondary Organic Aerosol Effects on the
15. Pulmonary Toxicity of Oxygen and Carbon Dioxide
17. Degraded amine. CO2 degraded amines overall showed only mild inflammato-
18. The results demonstrated that for different EPS products, such as lunch carriers, dishes and
19. The environment consequences from utilizing this technology have been
20. In aggregate, the results suggest that SOA causes mild cardiovascular responses that are dependant on composition. Research Funded by the Electric Power Research Institute.
estimated that students are inhaling six times more PCBS than they are receiving from dietary sources. However, LC-PCBs or metabolites have been rarely reported in the serum of population at risk. LC-PCBs are bioactivated to phenols and further to quinone electrophiles with genotoxic/carcinogenic potential. We hypothesized that phenolic LC-PCBs are subject to phase II conjugation and excretion via the urine. Our objective was to identify the final metabolites in the urine that could be potentially employed in the exposure analysis of LC-PCBs. Male Sprague Dawley rats (150-175 g) were housed in metabolism cages and received a single intraperitoneal injection of 600 μmol/kg body weight of PCB3. Control animals received corn oil only. Urine was collected every four hours, euthanized at 36 h, and serum was sampled at 40 h. The LC-MS analysis showed that PCB3 sulfates were the major metabolite in both urine and serum. Approximately 3% of the dose excreted in the urine as sulfate; with peak excretion occurring at 10-20 h post-exposure. The major metabolites were 4’PCB3sulfate, 3’PCB3sulfate, 2’PCB3sulfate and presumably a catechol sulfate. The whole urine was collected. The LC-MS analysis showed that PCB3 sulfates were the major metabolite in both urine and serum. Approximately 3% of the dose excreted in the urine as sulfate; with peak excretion occurring at 10-20 h post-exposure. The major metabolites were 4’PCB3sulfate, 3’PCB3sulfate, 2’PCB3sulfate and presumably a catechol sulfate. Serum level of 4’PCB3 was 0.18±0.16 μg/mL, while 4’OH-PCB3 was only 0.95±0.05 μg/mL. This is the first report that sulfated metabolites of PCBS are formed in vivo.


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The endogenous high affinity aryl hydrocarbon receptor agonist FICZ is formed upon irradiation of tryptophan with UV or visible light. To elucidate the mechanisms in more detail photooxidation products were analyzed by HPLC and LC/MS. The results revealed that the formation of FICZ was dependent on H2O2 since additions of catalase inhibited the reaction. It was also evident that indole-3-acetyladenine (I-3-A) was a precursor that spontaneously generated FICZ. Monoamine oxidases (MAOs) catalyze the oxidative deamination of monoamines resulting in the corresponding aldehydes. Human recombinant MAO A and MAO B were therefore incubated with tryptamine to yield I-3-A. After the reaction and a post treatment period the amounts of I-3-A and FICZ were measured. Both enzymes generated I-3-A, which was then spontaneously converted to FICZ. Subsequently, tryptophan was incubated with H2O2 at 37 °C in the dark. The mixture was concentrated on Sep-Pak C18, and HPLC and LC/MS analyses confirmed the formation of FICZ from tryptophan by H2O2. From these and other experiments it can be concluded that tryptophan can give rise to FICZ in an oxidative environment, i.e. containing or producing reactive oxygen species. In addition, when ever I-3-A is formed there is a possibility for spontaneous formation of FICZ. Altogether, this implies that there are several different conditions under which the endogenous Ah receptor ligand FICZ can be formed in the body. Of great interest in this respect is a recent demonstration of FICZ in the skin of vitiligo patients characterized by a massive epidermal oxidative stress and high levels of H2O2 (Shimizu et al., FASEB J. 2012). It therefore seems likely that FICZ is ubiquitously in the human body but expected to be present at low steady state levels under normal conditions.

1267 Significant Megaclass of Total Cancer Mortality among the US Non-White Population.


NCI and EPA collaborated to describe the U.S. population cancer mortality experience through the 1950s, 60s and 70’s. All mortality data are official statistics including 8 million records. For the 30 years, for all anatomic cancer sites, there was a significant difference in the percentage rate of increase between White and Non-white U.S. males. The White male increase was 17%. For the same period, Non-white males experienced a 46% increase. For both White and Non-white males highest rates of increase were in the Southeastern United States. The counties show up as streams along rivers. But the important unit of analysis is not the rivers, but U.S. Geological Survey (USGS) hydrologic units. Principal component analysis showed some USGS hydrologic units were more informative. USGS Region 08-02 St. Francis river valley experienced a 300% increase in White male lung cancer mortality in the 1970s, or a 10% increase per year. The heavily forested, rich bottom lands of the river valley were cut down and planted in cotton sprayed with DDTC. DDTC induces male rat lung cancers. The rates of high grade White male lung cancer counties lie predominately in the USGS Coastal Plains physiographic domain. During late Cenozoic and Tertiary Eras the Coastal Plains filled in with unconsolidated sands and sands, porous to pollutants leaching into shallow wells and aquifers. Relative risk of a red county being within the Coastal Plains was 5.3 times greater than being in another part of the U.S. Color coded counties showed a “dose-related” effect - the higher the county rate, the more likely it was located in the Coastal Plains. Authors solely responsible for conclusions herein and do not reflect authors’ institutions.

1268 Airborne Diacetyl from Cooking and Consumption of Microwave Popcorn: Estimation of Consumer Exposure with a Two-Zone Near-Field/Far-Field Model.

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Diacetyl is a volatile diketone used to impart a butter flavor to foods such as microwave popcorn. Occupational exposure to airborne diacetyl has been reported to be associated with obstructive lung disease, and recent concerns have emerged regarding exposures to diacetyl by consumers. Given the absence of data on the airborne concentrations in the home, we estimated consumer exposure using a two-zone near-field/far-field model in which the near-field is a hemisphere above the microwave popcorn bag, and conducted Monte Carlo uncertainty and sensitivity analyses. A recent study revealed that an average of 778.9 μg diacetyl was emitted during the popping, opening, and 40 minutes following opening. As part of the input to the model, we estimated that diacetyl is released into the breathing zone over a 40 minute period (a typical consumption duration) comprised of popping (2.5 min), opening (15 sec), and consumption (37.25 min). Based on available information, we estimated that 98.9% of the total diacetyl emitted was released during opening. The popcorn and the bag were assumed to be an arm length away from the nose of the consumer and present for the duration of the scenario. The estimated mean airborne diacetyl concentration in the breathing zone from one bag of microwave popcorn was 0.0030 ppm, whereas the 5th and 95th percentile concentrations were 0.0018 and 0.0048 ppm, respectively. Assuming complete absorption following inhalation and a breathing rate associated with light activity (0.66 l/min), the estimated maximum total intake by inhalation per bag was 5.3 μg. By comparison, the daily intake associated with occupational exposure at the eight hour TWA and 15 minute STEL Threshold Limit Values set by ACGIH (0.01 and 0.02 ppm, respectively) are 66 and four-fold greater, respectively.
The aryl hydrocarbon receptor (AHR) is a ligand activated transcription factor implicated in the regulation of diverse cellular processes. Previous studies in head and neck squamous cell carcinoma cell lines (HNSCC) have revealed considerable constitutive and ligand inducible AHR transcriptional activity. Antagonism of AHR activity through AHR antagonist treatment was found to greatly attenuate the highly metastatic and proliferative phenotype of these cells, suggesting that AHR plays a significant role in contributing to the ‘aggressive’ phenotype of these cells. Therefore, this study sought to use expression profiling to identify putative novel targets of the AHR repressed by AHR antagonist treatment that are associated with cancer cell survival and tumor invasiveness. Three growth factor targets were considered: amphiregulin (AREG), epiregulin (EREG) and platelet-derived growth factor A (PDGFA) identified by expression profiling and from previous studies. Quantitative PCR analysis revealed an attenuation of basal and/or induced gene expression levels of these growth factors in two HNSCC lines after AHR antagonist treatment. ELISA analysis revealed attenuation of both basal and induced protein levels of the growth factors examined in these cell lines. In addition, siRNA-mediated knock down of AHR exhibited attenuation of growth factor expression. In silico analysis revealed these growth factors possess dioxin-like response elements. Therefore, AREG, EREG and PDGFA were identified as three key growth factor targets of the AHR associated with metastatic phenotype of head and neck cancers.

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Naturally occurring flavonoids have been reported to possess paradoxical biological activities in vitro and in rodents; e.g. chemotherapy and exacerbation of estrogen-induced tumorgenesis. Previously, quercetin, kaempferol, and tamarixetin were reported to activate human PXR (hPXR). However, it is not known whether there is a structure activity relationship in the activation of hPXR by flavonoids and whether they activate human VDR (hVDR) and human GR (hGR). In the present study, we compared the effects of structurally related flavonoids (i.e. flavonol, galangin, daidzein, kaempferol, morin, quercetin, isorhamnetin, tamarixetin, myricetin, and syringetin) on the activity of hPXR, hVDR, and hGR. Only flavonol, galangin, isorhamnetin, tamarixetin, and quercetin activated hPXR, whereas none of them activated hGR or hVDR, as determined in dual-lucifase reporter gene assays in transfected HepG2 human hepatoma cells. Dose-response experiments indicated that the minimum effective concentration for hPXR activation by these flavonoids was 10-30 μM. Flavonol and galangin, but not isorhamnetin, tamarixetin, or quercetin, recruited steroid receptor coactivator-1 (SRC-1), SRC-2, and SRC-3 to the ligand-binding domain of hPXR, as assessed by mammalian two-hybrid assays. Flavonol and galangin bound to the ligand-binding domain of hPXR in a time-resolved fluorescence resonance energy transfer competitive ligand-binding assay. Overall, flavonoids activate hPXR in a receptor-selective and chemical-dependent manner. The addition of OH or OCH3 group at C2 and C5 positions in Ring B of flavonol appears to be unfavorable. Therefore, a minor change in chemical structure leads to abolishment of hPXR activation, implying that flavonoids may have differential impact on hPXR-mediated effects. [Supported by CHTR and MSFHR]

1278 Comparison of Serum Chemistry and Tissue Phenotypes in Aryl Hydrocarbon Receptor Knockout Rats and Mice.

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An aryl hydrocarbon receptor knockout (AHR-KO) rat was generated on a Sprague-Dawley outbred background in order to investigate the mode-of-action of 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) tumorgenesis. Aside from its role as a xenobiotic receptor, there is evidence that the aryl hydrocarbon receptor (AHR) plays a critical role in the development of some tissues. Previous reports describe a variety of tissue abnormalities in the liver, heart and kidney in AHR-KO mice. We conducted a comparative study of gross tissue pathology, histology and serum chemistry in AHR-KO rats and mice, and corresponding wild-types at 1, 6 and 12 weeks of age (n = 5 / sex / genotype / species) in order to evaluate the role of the AHR in tissue development across species. Adult AHR-KO mice, but not AHR-KO rats, had alterations in many serum chemistry markers associated with compromised liver function as compared to wild-type. Similarly, in adult AHR-KO mice, decreased liver-to-body weight and increased kidney- and heart-to-body weight ratios were observed. Similar changes were not observed in AHR-KO rats, save increased kidney-to-body weight ratios in females. Hepatic developmental abnormalities including portal hypercellularity, biliary reduplication, hepatocellular vacuolization and increased extramedullary hematopoiesis were observed in 1 week old AHR-KO mice and gradually dissipated across 6 and 12 week time points. Adult AHR-KO mice also had patent ductus venous. None of these histological or gross morphological liver phenotypes were observed in AHR-KO rats. In contrast, AHR-KO rats had severe urinary tract pathology including hydrocystoureter, hydronephrosis with secondary renal pelvic dilation, epithelial hyperplasia and mineralization leading to degenerative and inflammatory changes in cortical nephrons. Renal pathology was not observed in AHR-KO mice. Overall, these data indicate that the endogenous role of AHR in tissue development differs significantly between rats and mice.
Rheumatoid arthritis (RA) is a chronic autoimmune disease. It is characterized primarily by proliferation of cells in the synovial lining such as fibroblast-like synoviocytes (FLS). Under non-RA settings, FLS are a highly differentiated unicellular cell type. However, under inflammatory milieu, they become hyperplastic, invasive and highly migratory. Epidemiological studies have identified a positive correlation between cigarette smoking, a source of agonistic AHR ligands, and the progressive phenotype of RA. Thus, under inflammatory conditions, we hypothesize the AHR plays an important role in the expression of several growth factors in FLS. Our AHR gene knockdown data in IL-1β activated primary FLS suggest a positive link between ablation of AHR protein activity and suppression of inducibility of vascular endothelial growth factor-A (VEGF-A) and epiregulin (EREG) mRNA levels. GNF351, a potent AHR antagonist, was also shown to suppress cytokine-mediated upregulation of VEGF-A, EREG and basic fibroblast growth factor (FGF-2) expression, further validating the AHR dependency. ELISA was performed on supernatants collected from FLS-RA cells pre-exposed to GNF351, followed by stimulation with IL-1β, which revealed AHR antagonist-mediated suppression of VEGF-A activity. Cytokine dependent activation of growth factors has been shown to enhance FLS cell proliferation, which was attenuated by GNF351 pretreatment. Upon activation by cytokines, FLS have been shown to become invasive and highly migratory. Treatment of FLS with GNF351 mitigated cytokine-mediated expression of matrix metalloproteinases (MMP)-2 and -9 mRNA levels and also diminished FLS migratory. Treatment of FLS with GNF351 mitigated cytokine-mediated expression of matrix metalloproteinases (MMP)-2 and -9 mRNA levels and also diminished FLS migratory. Overall, these results suggest that the AHR may be a viable therapeutic target in amelioration of disease progression in rheumatoid arthritis.

We addressed responses of developing zebrafish to potential agonists for the pregnane X receptor (PXR); also Steroid Xenobiotic Receptor (SXR) and the aryl hydrocarbon receptor (AHR) to determine involvement and interaction of PXR and AHR signaling pathways in regulation of genes in cytochrome P450 subfamilies CYP2 and CYP3. Zebrafish embryos were exposed to 5-pregnen-3β-ol-20-one (pregnenolone; 1-10 μM) or to carrier (DMSO) for 24 h, starting at 48 hours post-fertilization (hpf), and then harvested at day 3. We also exposed one-day-old embryos to 3,3′,4,4′-4′-pentachlorobiphenyl (PCB126) or DMSO for 24 h and then held in clean water until day 4. Expression of PXR and selected CYP genes that are potential targets in the PXR signaling was assayed. Pregnenolone caused a concentration-dependent increase in mRNA expression of PXR, some CYP2AAs, CYP3A65 and CYP3C1, all of which peaked at 3 μM and then declined. An AHR agonist PCB126 also upregulated the transcript levels of those genes in most cases in a concentration-dependent manner. We next sought to examine the role of the PXR and AHR in the modified expression of those potential target genes by using morpholino antisense oligonucleotides (MO or morpholino) to block initiation of transcription of the PXR or AHR. Treatment of embryos with the PXR-MO, but not the control morpholino, partially inhibited pregnenolone-induced expression of PXR, CYP2 and CYP3 genes. Similarly, AHR2-MO treatment blocked PCB126-induced transcript expression of PXR and some CYP2 and CYP3 genes. The present study shows that PXR is not only self-upregulated, but also upregulated via activation of AHR2 in developing zebrafish. Selected zebrafish CYP2 and CYP3 genes appear to be in part under the regulation of PXR and AHR2. They include some CYP2AAs that were first identified in zebrafish. [Support: JSPS Postdoctoral Fellowships for Research Abroad no. 820 (A.K.), and NIH Superfund Research Program grant P42ES00738 (J.S.3)]
Inhibition of Heat Shock Protein-90 Prevents the Transactivation and Translocation of Human Constitutive ANDROSTANE RECEPTOR.


17-(Dimethylaminoethylamino)-17-demethoxygeldanamycin (17-DMAG) is currently in clinical trials for cancer treatment. 17-DMAG is a hydrophilic derivative of geldanamycin (GA), and its analog, 17-(allylamo)-17-demethoxygeldanamycin (17-AGA), all inhibitors of the molecular chaperone, heat shock protein-90 (HSP90). 17-DMAG offers a potential advantage over 17-AGA and GA due to its increased bioavailability, limited metabolism and reduced in vivo toxicity. HSP90 inhibition results in defective protein folding, conformation and assembly of its client proteins, promoting their destabilization and degradation. Client proteins of HSP90 include transmembrane tyrosine kinases (e.g., HER2, EGFR), intermediary signaling kinases (e.g., AKT, p53) and nuclear receptor family members (e.g., AHR, PXR). Previous reports demonstrated that HSP90 is a cytosolic tethering partner of the constitutive androstane receptor (CAR). The present investigation evaluated the potential effects of 17-DMAG on human CAR transcriptional activation and nuclear translocation. 17-DMAG treatments repressed CAR-mediated CYP450 induction in cultured human primary hepatocytes. Cell-based gene reporter assays verified that 17-DMGA inhibited hCAR in a dose-dependent manner. Intracellular localization studies revealed that phenobarbital-stimulated nuclear translocation of hCAR in hepatocytes was completely blocked by 17-DMAG, as well as GA. Immunoprecipitation analyses indicated that the suppression of hCAR activity by 17-DMAG resulted from an enhanced interaction between hCAR and HSP90. Mammalian two-hybrid assays further demonstrated that 17-DMAG disrupted the interaction of hCAR with the nuclear co-activator, SRC1. Together, the data indicate that these targeted HSP90 inhibitors markedly disrupt normal hCAR function, suggesting that their clinical use may potentially impact CAR-mediated drug metabolism and clearance.

Activation of Transient Receptor Potential Ankyrin-1 by Wood Smoke Particulate Material.

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Exposure to wood smoke particulate matter (WSMP) has been linked to exacerbation of asthma, development of causes chronic obstructive pulmonary disease (COPD), and premature death. Combustion-derived PM (cdPM) such as cigarette smoke (CS), diesel exhaust (DEP), and WSMP, activate transient receptor potential ankyrin-1 (TRPA1) which promotes neurogenic inflammation/redma and airway irritation/cough. The mechanism of TRPA1 activation by DEP and CS involves the electrophilic oxidant (3CK) and menthol-binding (ST) sites, and a novel mechanosensitive site. We hypothesized that WSMP would activate TRPA1 through one or more of these sites similar to other cdPM. Pine and mesquite PM were generated in the laboratory. Both types of WSMP particles activated TRPA1 in human TRPA1 over-expressing HEK-293 and primary mouse trigeminal (TG) neurons. WSMP also activated TRPA1 in A549, a human alveolar adenocarcinoma cell line, which has recently been shown to express TRPA1. HC-030031, a TRPA1 specific antagonist, attenuated the calcium flux due to WSMP treatment in both human A549 cells and mouse primary TG neurons. Several known chemical components of WSMP, including 3,5-diterbutylphenol and agastic acid were TRPA1 agonists. WSMP and agastic acid activated TRPA1 primarily via binding the 3CK site, based on inhibition of calcium flux by glutathione and mutation of the 3CK site. Conversely, 3,5-diterbutylphenol activated TRPA1 through the ST site. This study established the mechanisms by which WSMP and associated chemical components activated TRPA1 which may help tailor effective therapeutic treatments for WSMP pneumotoxicity. Support: NIEHS ES017431 and the University of Utah Undergraduate Research Opportunities Program.

Exploratory Phenotype and Molecular Analysis Study in ROR-Gamma Knockout Mice.


Retinoid-related orphan receptor g (RORg) is an orphan nuclear hormone receptor that is widely expressed in several tissues including liver and skeletal muscle. Previous gene expression studies knocking out the RORg knockout (KO) mice suggested a possible role for RORg in metabolism of steroids, bile acids, and xenobiotics (Jetten et al. 2007). RORg has also been shown to regulate several metabolism genes in skeletal muscle cells. The current study was designed to investigate the effects on pathologic and genetics of knocking out the RORg nuclear receptor in mice with a focus on liver and skeletal muscle tissues. Male and female wild type, heterozygote, or homozygote RORg KO mice (10/group) were assessed at 10 weeks of age. Study endpoints included clinical observations, body weights, organ weights, gross and microscopic examination of tissues, and gene expression evaluation using RNA isolated from liver, skeletal muscle, and spleen. There were no im pact with respect to clinical observations, body weight, and pathology on the phenotype of the heterozygous animals when compared with controls. Although no differences in clinical observations or body weights were observed in the homozygous animals, significant microscopic lesions were present in the thymus (thymic hyperplasia and lymphoma), spleen (increased cellularity), and lymph node (ab sent). These findings were not unexpected and are in line with established literature (Eberl and Litman 2003; Ueda et al. 2002). Pathway analysis of gene expression changes indicated that a majority of the altered pathways were related to immune response, an expected outcome. Skeletal muscle gene changes suggested a possible role for RORg in energy homeostasis. However, no perturbations in pathways related to toxicological consequences were noted. Taken together, the data showed that knocking out of RORg in mice led to expected changes in immune-associated pathology and gene expression, but did not reveal any additional phenotypic or genotypic toxicologically relevant effects.

Estrogen Receptor α and Aromatase Hydrocarbon Receptor Are Not the Sole Determinants of Aromatase-Mediated Growth Inhibition.

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Numerous studies have suggested that the aromatase hydrocarbon receptor (AhR) may be a potential therapeutic target for several human diseases, including estrogen receptor alpha (ERα) positive breast cancer. Aromatase (AF), an activator of AR signaling that is structurally related to phytochemicals called flavonoids, is currently in clinical trials for the treatment of solid tumors. It has been hypothesized that ERα may be a predictor for response to AF-mediated growth inhibition, with ERα-positive cell lines being sensitive to the drug, and ERα-negative cell lines being resistant. However, the scope of cell lines tested in the literature is limited. Here, we show that ERα-negative human breast cancer cell lines express greatly different amounts of AhR protein compared to ERα-positive human breast cancer cell lines, giving reason to further characterize AF’s efficacy in these models. Using cell counting assays to measure GFP values, we show that two ERα-negative breast cancer cell lines, MDA-MB-468 and Cal51, exhibit sensitivity to AF (GFP <1μM), suggesting that Erα is not the sole predictor of responsiveness to AF. To examine the mechanism of AF-mediated growth inhibition, a doxycycline-inducible AhR knockdown system was created in these two cell lines. We show that there is no significant difference in growth inhibition caused by AF treatment in cells where AhR is knocked down and cells with endogenous levels of AhR. This suggests that a portion of the mechanism involved in AF-mediated growth inhibition in MDA-MB-468 and Cal51 cells is AhR-independent, or that compensatory mechanisms may exist. Overall, this work suggests that while Erα is not the sole determinant of sensitivity to AF, this compound may also be inhibiting cell growth in an AhR-independent fashion. Supported by NIEHS Predoctoral Training Grant T32 ES007015.

Understanding the Physiological Role of Aromatase Hydrocarbon Receptor Repressor (AhRR) Using Gene Knock-Down and Targeted Mutagenesis in Zebrafish.

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The aromatase hydrocarbon receptor repressor (AhRR) is a transcriptional repressor of the aromatase hydrocarbon receptor (AhR) and is regulated by an AhR-dependent mechanism. In humans, AhR has been linked to tumorigenesis and reproductive defects. Zebrafish have two AhR genes; AhRRα regulates constitutive AHR signaling during development, while AhRRβ regulates polyaromatic hydrocarbon-induced gene expression. The goal of this study was to elucidate the endogenous role of AhRR in zebrafish development using two loss-of-function approaches: morpholino oligonucleotide (MO) knock-down and zinc finger nuclease (ZFN)-based gene targeting (knock-out). Zebrafish embryos were microinjected with MOs against AHRRα or AHRRβ to knock down the expression of these proteins, followed by microarray analysis of gene expression in embryos at 72 hours post-fertilization. In AHRRα-MO injected embryos, 97 genes were upregulated and 182 genes were downregulated. The latter included cone type photoreceptor-specific
genes, suggesting a role for AHRRs in photoreceptor development. In addition, AHRRs knock-down induced upregulation of embryonic hematopoietic (AhRR), suggesting a role in hematopoiesis. Knock-down of Ahrrs caused upregulation of 31 genes and downregulation of 85 genes, without enrichment of genes related to any specific biological process. These results suggest that AHRRs plays an important endogenous role in development. To understand the role of AHRRs beyond development, we targeted ahrr using ZFNs. Microinjection of ZFN mRNA into zebrafish embryos caused a 7 base pair frame shift mutation at the ahrr locus. We screened for germline mutants and identified a founder fish that was outcrossed to generate heterozygous mutant offspring. Heterozygous mutants were crossed to generate ahrr mutant homozygotes. Using these homozygotes we are investigating the role of AHRRs in normal physiology and toxicology. [Supported by NIH R01ES006272]

1289 Tissue-Specific Expression of Cytokines and Chemokines in TCDD-Treated B6 and Aryl Hydrocarbon Receptor Repressor Transgenic Mice.
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The aryl hydrocarbon receptor (AhR) has been identified as an important transcription factor regulating the responses of immune cells including T cells and dendritic cells but the role of the AhR Repressor (AhRR) is less clear. The AhRR has been first described to suppress AhR activity as an inducible negative feedback loop, then again there are many unanswered questions, as we know now that the basal or induced cytokrome P450 1A1 (CYP1A1) mRNA level rarely correlates with AhRR expression. To gain insight into the function of AhRR in vivo we developed a transgenic AhRR B6 mouse (AhRR Tg). These transgenic mice express significantly higher levels of AhRR mRNA in all tissues examined such as liver, lymphnode, kidney, spleen, and thymus. Treatment of B6 and AhRR Tg mice with TCDD, a single dose of 1 μg/kg body weight, or with corn-oil (vehicle) given 5 days after treatment with TCDD from our own breeding five days after treatment with TCDD, male wild-type (wt) and AhR-deficient mice (originating from Finish Public Health Institute, Kuopio, Finland) from our own breeding five days after treatment with TCDD, single dose, 25 μg/kg body weight, or with corn-oil (vehicle) given by oral gavage. Subsequently, RNA was isolated and Microarray analysis was performed using Agilent Whole Mouse Genome Oligo Microarray 4x44K. In wt mice, 282 genes were more than 2-fold up-regulated by TCDD in the liver including some of well-known AhR target genes like cytokrome P450 (CYP) 1A1, 1A2 and 1B1; 152 genes were up-regulated in the kidney. In AhR-deficient mice, 242 hepatic and 90 renal genes showed an increased expression after TCDD treatment. Among those, 15 hepatic and 13 renal genes were up-regulated in both wt and AhR-deficient animals. None of the core genes of the AhR gene battery except CYP1B1 was up-regulated in AhR-deficient mice. The AhR-independent, TCDD-regulated genes comprise genes involved in lipid, carbohydrate, and amino acid metabolism, and in immune regulation. In wt mice, 203 hepatic and 20 renal genes were more than 2-fold down-regulated. In AhR-deficient mice, 234 hepatic genes including CYP1A2 and CYP3A4 and 235 renal genes showed a decreased expression. In addition, 18 hepatic genes were down-regulated in both wt and AhR-deficient animals. Our results might provide new insight into the role of the AhR in the immune system.

1290 Caloric Restriction Induces Nrf2-Dependent Effects on Lipid Metabolism.
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Toxascrambling has identified the Nrf2 transcriptional pathway as being highly constitutive to chemical promiscuity. Nrf2 is classically defined to have a role in mediating cytoprotection against oxidative stress, but recent work points to a more fundamental metabolic role in lipid metabolism. Studies have indicated that the Nrf2/Keap1 pathway is inducible in adipose tissue, and Nrf2 may have a role in the fundamental metabolic role in lipid metabolism. Studies have indicated that the diating cytoprotection against oxidative stress, but recent work points to a more...
1293 Coexposure to Aryl Hydrocarbon Receptor and Thyroid Receptor Agonists Enhances Induction of Responsive Genes in Cultured Frog Cells and Prometamorphic Frogs.

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Amphibian metamorphosis is a postembryonic developmental process driven by thyroid hormone (TH) and mediated by the thyroid receptor (TR). 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) can disrupt TH function in humans and other species. Toxicity of dioxin-like compounds is mediated by the aryl hydrocarbon receptor (AhR). Here we used the African Clawed Frog (Xenopus laevis) as a model to probe the interaction of TR and AhR signaling. We quantified relative mRNA expression of both TR and AhR target genes in prometamorphic tadpoles and in XLK-WG cells following exposure to both TR and AhR agonists. Tadpoles (NF 53-55) were exposed to 5 nM TCDD and/or 10 nM T3 for 18 hours. TR targets (βE2R, βRbα, and TRβb) were not strongly induced by TCDD alone, and AhR target (CYP1A6) was not responsive to T3. However, exposure to both compounds resulted in elevated transcriptional responses of both transcript classes. Compared with T3-exposed animals, βE2R, βRbα, and TRβb mRNAs were 40-70% more abundant following co-exposure with TCDD. Similarly, CYP1A6 mRNA was ~50% more abundant in tadpoles following T3 co-treatment than with exposure to TCDD only. Comparable patterns were observed in XLK-WG cells. The effect of TCDD on circulating TH varies with species and can differ for T4 and T3. While the superinduction of TH-responsive frog genes could involve TCDD-related alteration of TH levels, the requirement for co-treatment and the effect in cultured cells suggest the possibility of a transcriptional mechanism. We examined the promoter regions of βE2R, TR-βb, and CYP1A6, finding potential cognate binding elements for each receptor. Overall, our results suggest some degree of co-regulation of the expression of several genes by both AhR and TR. Frogs display great insensitivity to the induction of TR-regulated genes by AHR may represent an additional mechanism to protect them from xenobiotic or endogenous AHR agonists. (NIH; R15 ES011130)

1294 A Global Genomic Screening Strategy Reveals Diverse Activators of Constitutive Activated Receptor (CAR).

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A comprehensive survey of conditions that activate CAR in the mouse liver has not been carried out but would be useful in understanding their impact on CAR-dependent liver tumor induction. A gene signature dependent on CAR activation was identified by comparing the transcript profiles after exposure to three CAR activators (phenoxybenzal, TCPOBO, CFCO) in wild-type and CAR-null mice. In independent experiments using transcript profiles from the livers of chemically-exposed male or female mice, the signature correctly predicted activation of 3 CAR activators but not 9 activators of other pathways. The signature was used with 5 classification methods (e.g., support vector machines, K-nearest neighbors) to identify conditions in which CAR was activated in an Affymetrix compendium of ~750 mouse liver transcript comparisons encompassing a broad range of chemical, dietary and genetic perturbations. We found that CAR is activated by a large number of chemicals, dietary regimens and genetic mutations. Specific and novel findings include activation by i) 2 PXR activators in a PXR-dependent manner indicating crosstalk between CAR and PXR, ii) 12 out of 15 chemical and triglyceride activators of PPARα to greater levels in PPARα-null mice than in wild type mice indicating that most PPARα activators are also CAR activators and there exists antagonism between CAR and PPARα, and iii) null mutations in a number of transcription factors (AhR, Fxr, Hnf1a, Pxr) that control expression of genes involved in metabolism of exogenous and endogenous chemicals. The findings increase our understanding of the factors that impact CAR activation and that could contribute to increases in CAR-dependent liver tumors. This abstract does not represent EPA policy.

1295 Effects of Munitions Compounds on Xenobiotic-Activated Nuclear Receptors and Cell Signaling Pathways.

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Exposure to certain munitions compounds is known to alter physiological functions in test organisms, however little is known about their molecular and cellular effects. The objective of this study to characterize the effects of new and existing munitions compounds on xenobiotic-activated nuclear receptors and cell signaling pathways. Munitions compounds (0.01-10 mg/l) (e.g., nitroaromatics, cyclic nitramines, and new insensitive munitions) were added to wells containing transfected cells containing high constitutive levels of nuclear receptors for 24 h, then examined for nuclear activation according to in-house and commercially available kits. In general, nitroaromatic compounds had greater effects on nuclear receptor activity decreased nuclear receptor activity than other munitions compounds. Nitroaromatic munitions decreased activation of constitutive androstane receptor (CAR), peroxisomal proliferator activated receptors (PPAR), liver X receptor (LXR) at higher concentrations examined, whereas some nitroaromatic compounds activated the Ah receptor. Effects on 45 cell signaling pathways were examined using the Signal Finder multi-pathway reporter array kit that reverse-transfected transcription factor-firefly luciferase reporter plasmids into HepaRG liver cells. Cells were then exposed to munitions compounds for 24 h, then examined for signal pathway activation. Trinitrotoluene (TNT) and its environmental byproducts differentially activated cell signaling pathways, with at least two pathways commonly shared (AhR and Nrf2/Nrf1). 2,4-Dinitroanisole, a new TNT replacement munition, activated fewer cell signaling pathways than TNT. These results create a more comprehensive picture of munitions effects within cells and potential downstream effects within organisms. Furthermore, rapid natural bioassays, such as the ones described above, may help material scientist predict biochemical pathways that may be impacted by newly developed munitions earlier in the developmental process.

1296 The Aryl Hydrocarbon Receptor Depletion in Human MDA-MB-231 Breast Cancer Cell Line Attenuates In Vivo Tumor Growth and Pulmonary Metastasis in a Nude Mouse Model.

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The aryl hydrocarbon receptor (AhR) is a ligand activated transcription factor that is best characterized for its role in mediating the toxic responses elicited by environmental poly aromatic hydrocarbons (PAH). However there is compelling evidence for a PAH-independent role of AhR in breast cancer. AhR is overexpressed and constitutively activated in several human breast carcinoma cell lines and shows strong correlation with the degree of the tumor malignancy. In the present study we examine the effect of depleting AhR expression in the highly metastatic MDA-MB-231 human breast cancer cell line on tumor growth and experimental pulmonary metastasis in a nude mouse model. The AhR was stably knocked down using shRNA targeting AhR gene. Clonal cell line of MD231 HBC with approximately 80% depletion of AhR and its scramble control cells were transfected to stably and constitutively express luciferase gene for the purpose of in vivo bioluminescence imaging of tumors. Cells were injected either orthotopically in the mammary fat pad for monitoring the primary tumor growth, or intravenously in the lateral tail vein, for monitoring pulmonary experimental metastasis. Results showed that knock down (KD) of AhR expression significantly reduced incidence of mice with tumor growth, prolonged the latency and reduced tumor size in those mice which developed tumors compared to controls. Mice with AhR KD in the experimental metastasis group showed a significant reduction in the number of animals with lung metastasis compared to control, and decreased numbers and sizes of pulmonary metastasis in those, which manifested metastasis. Global gene expression profiling identified sets of genes associated with tumor growth and metastasis that have been subsequently altered following depleting of AhR, and are likely to account for these phenotypes. Our data provide the first in vivo evidence for the role of AhR in regulating breast cancer metastasis and identified it as a therapeutic target for metastatic breast cancer.

1297 Species Specificity of the Peroxisome Proliferator Response in Primary Hepatocytes.

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Perturbations induced by environmental chemicals often lead to different physiological outcomes in humans and rodents, calling into question the applicability of animal-based toxicity testing for human risk assessment. For example, peroxisome proliferators, which include several purported endogenous fatty acids as well as a number of synthetic ligands, induce lipid metabolism enzymes in humans and have been used successfully as therapeutic strategies for various dyslipidemias and diabetes. For rodents, in addition to their role in regulating fatty acid metabolism, peroxisome proliferators also influence peroxisome assembly, inflammatory responses,
and cellular proliferation. Understanding the basis for differences in clinical repercussions across species is essential to translating information from animal models to human health.

Here we present a genomic characterization of the response of human and rat primary hepatocytes to the PPARα-specific agonist GW7647. We used gene expression microarray experiments to compile a highly resolved time- and dose-response characterization of the differences in the PPARα-mediated response in these two species. Consistent with the qualitative differences in the response to peroxisome proliferators, we identified a substantially larger set of differentially expressed genes in rat cells compared to human. These additional genes include a suite of developmental processes that may account for the increased toxicity of peroxisome proliferator agents in rodents. Surprisingly, in both species, we found that the canonical nuclear receptor response mechanism accounts for only a small fraction of detected response, suggesting a larger role for non-genomic mechanisms than was previously thought.

1298 Development of a Rapid, Cell-Based Screen for Disruptors of the Retinol (Vitamin A) Signaling Pathway.

A central goal in the 2007 National Research Council’s vision for Toxicity Testing in the 21st Century is the development of a comprehensive suite of in vitro tests covering the major signaling pathways (toxicity pathways) in mammals that can be used for evaluating chemical toxicity while reducing or eliminating the use of animals. Our research focuses on the development of in vitro screens for detecting chemicals in food addittives, supplements, and contaminants that can disrupt the major cellular signaling pathways in mammals. One of these pathways, the retinol (vitamin A) signaling pathway, is essential for life in all mammals. It is required for both normal embryonic development and maintenance of cellular phenotype in adult organisms; chemicals that cause even minor interference with its normal function and output are potential fetal and adult toxicants. We have developed a rapid (24 h) screen for detecting chemicals that disrupt this essential signaling pathway. It uses the mouse pluripotent P19 stem cell line and a medium-throughput 96-well format gene-expression assay to detect disrupting chemicals. It has detected all known retinoid signaling pathway disruptors and some chemicals that are closely related, structurally, to known disruptors. Significantly, it also has detected members of a class of chemicals, the endocrine disruptors, which have not been associated previously with disruption of this pathway; chemicals that caused disruption included xenoestrogens (DES, BPA, 4-nonylphenol, and genistein and phthalate esters (dibutyl phthalate and diphenyl phthalate but not bis(2-ethylhexyl) phthalate). The effects of members of this class of chemicals on the pathway suggest the existence of an additional non-genomic mechanism of action by which some endocrine disrupting chemicals cause toxicity.

1299 Asian Ginseng (Panax ginseng) Potentiates Ethanol-Induced Cardiovascular Dysfunction in Medaka Embryogenesis (Oryzias latipes).
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Alcohol is a teratogen, induces fetal alcohol spectrum disorder (FASD) which has serious central nervous system (CNS), cardiovascular, and craniofacial defects affecting the entire lifetime of an individual. Prevention of FASD, other than women abstaining from drinking alcohol during pregnancy, is not known. The synthetic drugs recommended for the treatment of alcoholism cannot be used by women during pregnancy which led us to investigate on natural products. Due to ethical constraints FASD studies in humans are very limited and several animal models are used to understand the molecular mechanisms. We have observed that development of ethanol-induced teratogenesis in medaka (Oryzias latipes) embryos genotypes which are analogous to human FASD phenotypes. We hypothesize that ethanol metabolism generates oxidative stress which can disrupt embryonic development of medaka. In the present experiment, we have used root extracts of Asian ginseng (Panax ginseng) as a preventive agent of FASD. Fertilized medaka eggs within 4 h post fertilization (hpf) were exposed to methanolic extracts (50–100 µg/ml) of ginseng root (PG) or ethanol (300 mM) either alone or in combination. After 48 h of treatment the viable embryos were transferred to clean hatching solution and on 6 dpf the embryos were examined for vessel circulation followed by mRNA analyses of enzymes related to ethanol metabolism and oxidative stress. It was observed that ethanol (300 mM) alone was able to disrupt vessel circulation and treatment of PG (50–100 µg/ml) with ethanol was able to enhance the effect; PG (100 µg/ml) alone has no effect. mRNA analysis of alcohol metabolizing enzymes or oxidative stress-related enzymes did not show any significant alterations in any of these treatment conditions. It was concluded that potentiation of ethanol-induced cardiovascular deformities in medaka by PG may be mediated through a different mechanism rather than oxidative stress.

1300 Toxicology Profile of Virginia Cedarwood Oil Delivered via Dermal Application.
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Virginia cedarwood oil (CWO), which is extracted from Juniperus virginiana trees, is widely used as an insect repellent and a fragrance in cosmetic formulations resulting in human exposure. To investigate the toxicological effects of CWO, 90-day studies in F344 rats and B6C3F1/N mice were conducted. CWO was administered by dermal application to male and female rats at concentrations of 0, 6.25, 12.5, 25, 50% (v/v) in 95% ethanol or 100% daily (excluding weekends) for up to 14 weeks with an untreated control included. In rats, there were no effects on survival or body weights except in the 100% males in which 2 rats were terminated on day 79 due to severe skin lesions and a 13% lower mean body weight than that of the untreated control. All male and female rats in the 100% group were terminated during week 10 due to the severity of skin lesions. Final group mean body weights for females given 12.5, 25 or 50% and males given 50% were less than 90% of the controls. At the site of application, treatment related irritation, thickened skin, and ulcer formation were observed in all treated groups of rats and mice, except the 6.25% groups in rats. Clinical pathology showed an inflammatory leukogram secondary to dermal ulcers/inflammation in both rats and mice; mice also had a decreased erythrocyt. In rats and mice there were dose-dependent increases in incidences and severity of skin lesions at the site of application, and bone marrow hyperplasia. In rats, there were dose-dependent increases in incidences and severity of kidney lesions in males. Male mice demonstrated increased incidences of kidney nephropathy and thymus atrophy.

1301 Novel Antiplatelet Activity of Protocatechuic Acid through Inhibition of High Shear Stress-Induced Platelet Aggregation.
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Bleeding is the common and serious adverse effect of currently available antiplatelet drugs. Many efforts are being made to develop novel anti-thrombotic agents without bleeding risks. Shear stress-induced platelet aggregation (SIPA) which occurs under abnormally high shear stress plays a crucial role in the development of arterial thrombotic diseases. Here we demonstrated that protocatechuic acid (PCA), a bioactive phytochemical from Lonicera (honeysuckle) flowers, selectively and potently inhibits high shear (>10,000 s−1)-induced platelet aggregation. In isolated human platelets, PCA decreased SIPA and attenuated accompanying platelet activation including intracellular calcium mobilization, granule secretion and adhesion receptor expression. Anti-SIPA effect of PCA was mediated through blockade of von Willebrand factor (vWF) binding to activated glycoprotein IIb/IIIa; a primary and initial event for the accomplishment of SIPA. Consipicuously, PCA did not inhibit platelet aggregation induced by other endogenous agonists like collagen, thrombin or ADP that are important both in pathological thrombosis and normal haemostasis. Anti-thrombotic effects of PCA were confirmed in vivo in rat arterial thrombosis where PCA significantly delayed the arterial occlusion induced by FeCl3. Of a particular note, PCA did not increase bleeding time in rat tail trans-section model, while conventional anti-platelet drugs, aspirin and clopidogrel substantially prolonged it. Collectively, these results suggest that PCA may be a novel anti-platelet agent which can prevent thrombosis without increasing bleeding risks.

1302 The Soy-Associated Phytoestrogen, Genistein, Does Not Protect Against Alcohol-Induced Osteoporosis in Male Mice.
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Alcohol abuse acts as a risk factor for osteoporosis by increasing osteoclast activity and decreasing osteoblast activity in bone. These effects can be reversed by estradiol. Soy diets are also suggested to have protective effects on bone loss in men and
women, as a result of the presence of soy protein-associated phytoestrogens such as genistein and daidzein. In this study, male PF pair fed (PF) a casein diet, an EtOH diet or EtOH diet supplemented with genistein (250 mg/kg) for 8 weeks. Ex vivo microCT analyses of formalin fixed tibia from each group revealed a significant decrease in trabecular bone in the EtOH group in comparison with the pair-fed control in regards to bone volume (BV/TV), trabecular number (Tb.N), and trabecular separation (Tb.Sp), (p<0.05). Post Hoc comparisons showed that EtOH diet or EtOH diet supplemented with genistein (250 mg/kg) for 8 weeks. Ex vivo microCT analyses of formalin fixed tibia from each group revealed a significant decrease in trabecular bone in the EtOH group in comparison with the pair-fed control in regards to bone volume (BV/TV), trabecular number (Tb.N), and trabecular separation (Tb.Sp). Interestingly, there was an increase in trabecular thickness (Tb.Th) in the PF+genistein group compared to the PF, (p<0.05), suggesting genistein can affect bone remodeling. In ex vivo bone marrow cultures, EtOH exposure decreased the number of pre-osteoblasts compared to PF controls. In contrast, exposure to EtOH+genistein increased pre-osteoblast numbers compared to the EtOH-treated group, (p<0.05). These findings suggest that genistein has a partial protective effect on bone formation. In conclusion, genistein does not protect against ethanol induced bone loss despite increasing osteoblastogenesis. Supported in part by R01 AA18282 (M.J.R.) and UAMS INBRE award 8 P20 GM103429-11.

1303 Toxicological Safety Assessment of a Hydroethanolic Extract of Caralluma fimbriata.

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A toxicological safety assessment was conducted on a hydroethanolic extract of Caralluma fimbriata, an ingredient marketed as an appetite suppressant, to predict safety with oral consumption by humans. Caralluma fimbriata extract (CFE) is standardized to contain no less than 25% pregnane glycosides and no less than 10% saponin glycosides. The extract is ≥98% soluble in water. CFE is currently sold as a trademarked ingredient Slimaluma™. Two genotoxicity studies were conducted and no evidence of mutagenicity or genotoxicity was observed in the presence of or absence of a rat liver S9 metabolic activation system at concentrations up to 5,000 µg of extract/ml in a chromosomal aberration assay. Studies conducted in Wistar rats included a 14-day acute oral toxicity study, and a 90-day repeated oral toxicity study. A 6-month repeated oral toxicity study was conducted in Sprague-Dawley rats. In the 14-day study, the NOAEL was determined to be 5 g/kg bw. While a few statistically significant (p<0.05) findings were observed in the 6-month study and 90-day study, these were considered to be a sound basis for conducting a 6-month study. In the 6-month Sprague-Dawley rat study, the observed effect level (NOEL) was concluded to be 1,000 mg/kg bw/d, the highest dose group tested. Finally, in a developmental toxicity study in Sprague-Dawley rats no fetal abnormalities related to administration of the test article were observed.

1304 The Antidiabetic Activities of the Soft Drink Leaf Extract of Phyllanthus amarus (Order Euphorbiaceae) in Laboratory Animals.

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Phyllanthus amarus Schum (Family Euphorbiaceae) is an annual herbal shrub which has been used in traditional medicine in Nigeria to treat some disease conditions. This study evaluated the soft drink extract (SDE) of the plant for anti-diabetic activities in rats. SDE was prepared by dissolving fresh aerial parts of the plant in 7up soft drink for 48 h, filtered, lyophilized and then used for the pharmacological investigations. Standard phytochemical methods were used to test for the presence of phytoactive compounds in the plant. Acute toxicity was carried out in mice to determine safe doses for this plant extract. The anti-diabetic activities of the SDE of the plant were assessed using some standard tests as well as histological changes in liver, kidney and pancreas. Diabetes mellitus was induced in rats using alloxan while glibenclamide at 0.2mg/kg was the reference drug used in this study. The SDE at 200 and 400mg/kg body weight caused a significant reduction of fasting blood glucose, significant change in the oral glucose tolerance test, marked effect in the hyperglycaemic activity test, and pronounced reduction on the glucose, cholesterol and triglyceride levels of diabetic rats. The haemograms of all treated diabetic rats experienced recovery. Histopathologically, the liver of the diabetic non-treated and glibenclamide-treated groups showed widespread vacuolar change in the hepatocytes but there was no visible lesion seen in the kidney and pancreas of extract-treated and glibenclamide-treated groups. No lesion was also seen in the liver of SDE-treated group.

In conclusion, the results from this study may have validated the traditional basis for the use of Phyllanthus amarus as antidiabetic agent. The pharmacological activities noted in this study may be attributed to the presence of flavonoids and other phenolics contained in this plant. At the doses used, SDE also appeared safer than glibenclamide even though the latter is more potent.

1305 Protective Effects of Methanolic Root Extract of Balanites aegyptiaca (L.) Delile on Carbon Tetrachloride (CCH)-Induced Hepatotoxicity in Rats.

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Objective: Balanites aegyptiaca (BA), a widely grown desert tree found in Africa and Asia is used as a remedy for several ailments including liver and spleen diseases. This study evaluated the protective activity of the methanol extract of BA root by direct (curative and prophylactic models) and indirect (barbiturate-induced sleep model) methods in CCH -induced hepatotoxic rats.

Methods: In the curative study, 0.5 ml/kg CCH4 was given intraperitoneally (ip) on alternate days, for 5 days, followed by 100, 200 and 400mg extract/kg orally daily, for 7 days. Other groups received the vehicle only, CCH4 and the vehicle, and CCH4 and the standard drug, silymarin (100 mg/kg/day) respectively. In the prophylactic study, the rats received the treatment respectively for the first 7 days followed by CCH4 as above. Degree of liver protection was measured in the rats by biochemical parameters (serum alanine transaminase, aspartate transaminase, alkaline phosphatase, albumin, bilirubin, triglycerides and cholesterol levels) monitoring, physical (weight) and histopathological changes in the liver. The indirect method was same as in prophylactic study but on the 3th day, the rats were given 25mg pentobarbitone sodium/kg ip and observed for onset and duration of sleep.

Results: Pre-treatment with the extract significantly (P<0.05) decreased CCH4 induced elevation of serum levels of the biochemical parameters, better than silymarin. In the curative study, its effect was comparable with that of silymarin. It also significantly decreased pentobarbitone sleeping time. There was improved tissue histopathology in both studies.

Conclusion: These results showed that BA root has significant hepatoprotective activities and could be useful as ‘lead’ in the development of new hepatoprotective agent.

1306 The Effects of Citrus Auraptene in Combination with All-Trans Retinoic Acid on Human Squamous Cell Carcinoma in a Xenograft Model.

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Over 2,000,000 new cases of non-melanoma skin cancer (NMSC) are expected in the United States in 2012. Our group has previously demonstrated that citrus auraptene (AUR), combined with all-trans retinoic acid (ATRA), can inhibit the growth of human skin squamous cell carcinoma (SCC) tumors from the cell line SRB12-p9 (PWIT) in SCID/bg mice. ATRA is known to inhibit the STAT3 pathway, whereas AUR suppresses the activation of NF-κB and induces antioxidative enzymes such as glutathione S-transferase (GST). NF-κB and STAT3 are both hyper-activated in cancer. We hypothesize that suppressing both pathways by these compounds will produce a greater chemopreventive effect than by using either agent alone. Groups of 10 each female SCID/bg mice were injected with 1 x 10^6 PWIT cells, s.c. Mice were dosed with AUR (0, 100, 200, 400 mg/kg bw) ± ATRA (400 mg/kg bw/day, beginning on day of tumor study injection and 4 days/wk for the duration of the study (48 days). Tumor volumes were estimated by caliper. At the end of the tumor study, necropsies were performed, and tissues taken for further analyses. Statistical analyses revealed no effect of AUR by itself on tumor volumes. ATRA alone suppressed tumor volume by 16-fold. However, ATRA was so effective at blocking tumor volume that any additional effects of AUR may have been masked. ATRA also caused a loss of body weight, but the liver/body weight ratios were not different across groups, and the activity index of the mice was within range. In conclusion, this study will be repeated using lower doses of ATRA, in combination with AUR to determine the minimum effective dose of ATRA, (known to be toxic) while assessing the effect of the AUR. (5R21CA149761-02)
Toxicological Evaluation of the Aqueous Extract of Morinda morindoides Root Bark.

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Toxicological evaluation of Morinda morindoides root bark was carried out using hematochemical and serum biochemical changes accompanying prolonged administration of the aqueous leaf extract of the plant in Wistar rats. Twenty rats were randomly but equally divided into four groups. Rats in groups I were administered with 0.9% physiological saline (10ml/kg b.w) as the control animals, while rats in groups II, III and IV were administered with aqueous extract at 400, 800 and 1600 mg/kg body weight respectively once daily for 28 days according to the acute toxicity study carried out. Parameters evaluated were complete blood count, serum protein, metabolites and enzymes. Samples from liver, spleen, heart, and kidney were processed for histopathology. Data were expressed as mean±Standard Error of Mean and analyzed using one way ANOVA. Difference of means are considered statistically significant at p<0.05.

There were no significant changes in the hematometrical parameters between the treatment groups and the control group. Rats in group IV exhibited significant increase in the serum levels of alkaline phosphatase, aspartate amino transaminase, alanine amino transferase and creatinine. Histopathology revealed mild diffuse degeneration of the liver, mild tubular nephropathy and mononuclear cellular infiltration in the heart. It was concluded that ethnomedicinal application of Morinda morindoides is quite safe at lower doses but it is hepatotoxic, nephrotoxic and cardiotoxic at higher doses.

Antidiarrheal Evaluation of Aqueous and Ethanolic Leaf Extracts of Acacia sieberiana DC. (Fabaceae) in Albino Rats.


Sequel to reports on the use of Acacia sieberiana leaves in the management of animal diarrhea in some parts of Plateau State Nigeria, the antidiarrheal activity of aqueous and ethanolic extracts of the leaves of A. sieberiana were evaluated in albino rats. Studies were carried out on castor oil induced diarrhea and gastrointestinal transit at doses of 300, 600, and 1200 mg/Kg body weight. Both extracts did not show any significant (P>0.05) inhibitory effect in both assays when compared to the negative controls. The standard drugs, loperamide (10 mg/Kg) and atropine sulphate (3 mg/Kg) significantly (P<0.05) inhibited diarrhea and reduced the distance travelled by the activated charcoal in the intestine respectively. Phytochemical screenings of the extracts revealed the presence of saponins, cardiac glycosides, and flavonoids in both extracts, while tannins were present only in the ethanolic extract.

In the toxicity test (limit test), no deaths or any apparent signs of toxicity or side effects were observed in the animals following the oral administration of both extracts at 2000 mg/Kg. The traditional use of A. sieberiana leaves in the management of diarrhea could not be justified by this study. Further work to assess the antibacterial, antiviral, anthelmintic and antiprotozoal activity of the leaves and antidiarrheal evaluation of other parts of the plant is necessary to verify the folkloric claims for its use in diarrheal management.

Protection by Fish Oil against D-Galactosamine-Induced Hepatic Injury and Biochemical Alterations in the Plasma and Liver of Rats.


The present study has investigated the actions of fish oil (FO) on D-galactosamine (GalN)-induced changes in plasma glucose (GLC), bilirubin, total proteins (TP) and enzymatic indices of hepatic damage; and in the levels of plasma and hepatic triglycerides (TG) and cholesterol (CHOL). Groups of 6 male Sprague-Dawley rats, 220-50 g in weight, were treated for 3 consecutive days with a 600 mg/kg oral dose of FO, as an emulsion in 0.9% NaCl-tween 20. On day 4, the rats received a single intraperitoneal, 400 mg/kg, dose of GalN in 0.9% NaCl. Additional rats received only the vehicles of the treatment solutions (control group), only GalN or only FO. All the rats were sacrificed by decapitation under isoflurane anesthesia 24 hr after receiving GalN, and their blood and livers were collected. The plasma fraction was analyzed for GLC, TG, CHOL, total (TB) and direct (DB) bilirubin, TP, alanine transaminase (ALT), aspartate transaminase (AST) and alkaline phosphatase (ALP). A portion (1.5 ml) from each plasma was homogenate in phosphate buffered saline pH 7.5 which was analyzed for its contents in TG and CHOL. Relative to control values, GalN increased the plasma GLC (41%), ALT (>200%), AST (>110%), ALP (+ 54%), TB (6.3-fold), DB (46-fold), TG (5.3-fold) and CHOL (1.6-fold) while decreasing those of TP (by 59%) (all at p<0.01). In addition, GalN increased the liver weight to body weight ratio (35%) and the levels of TG (61%) and CHOL (400%) significantly (p<0.01 vs. control). A pretreatment with FO was able to attenuate the changes in plasma and liver components as well as in the liver weight to body weight ratio to a significant extent (p<0.05). By itself, FO was found to raise the plasma (98%) and liver TG (23%) and the liver CHOL (35%) significantly (p<0.05), but not as much as GalN. In conclusion, FO is found to protect the liver against GalN-induced hepatic injury, dysfunction and fat buildup and to attenuate accompanying changes in circulating GLC, TP, TG and CHOL.

The Phototoxicity of Tagetes Oil and Absolute Evaluated Utilizing a Human Epidermal Model and Patch Testing.

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Tagetes essential oil and absolute (Tagetes minuta L.) are utilized as fragrance ingredients in a variety of consumer and personal care products. Spectral analysis shows that the materials absorb UV wavelengths in the critical region of 290 – 700nm, and therefore have the potential to be photoactivated. In order to determine the potential for phototoxic effects, both materials were assessed in a phototoxicity test on a three dimensional human epidermal model (EST1000). This assay is based upon a comparison of the cytotoxic effects of the test material following exposure to a non-toxic dose of UV light. Cytotoxicity is expressed as a reduction in the mitochondrial conversion of MTT [(3-4,5-dimethyl thiazole 2-yl) 2,5-diphenyl-tetrazoliumbromide] to formazan. A material is considered to have a phototoxic potential if one or more concentrations +UVA (60 min with 61/cm²) results in a decrease in cell viability of ≥30%, when compared to the identical concentration –UVA. Tagetes essential oil and absolute were tested at concentrations ranging from 0.1% to 10% (v/v) in 1:3 ethanol:Depot, in a 24-hour exposure period. Phototoxicity patch test was then conducted in a group of human volunteers, at a concentration identified to be non-phototoxic. In the EST1000 model, no phototoxicity was observed to Tagetes essential oil over the range of concentrations tested. However, within this same model, phototoxic effects were observed following exposure to Tagetes absolute at concentrations as low as 0.1%. Based on the dose response in these in vitro assays, a concentration of 0.01% was identified as being non-phototoxic. This concentration (0.01% in 1:3 EtOH:DEP) of Tagetes essential oil and absolute was confirmed as having no phototoxic effects in humans following the phototoxicity patch test.

Saponins from Platyodon grandiflorum and Its Constituent Platycein D Regulates Fatty Acid Metabolism via an AMP-Activated Protein Kinase Dependent Signaling Pathway.

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The saponins from the roots of Platyodon grandiflorum (CKS) have a variety of pharmacological properties, including antioxidant and hepatoprotective properties. This study was conducted to suggest the role of AMP-activated protein kinase (AMPK) pathway in the anti-obesity effect of CKS. We evaluated body weight, liver histology, and hepatic triglyceride content in high-fat diet (HFD)-fed mice treated with CKS. In addition, we characterized the underlying mechanism of CKS’s effects in HepG2 cells by Western blot and RT-PCR analysis. CKS administration attenuated fat accumulation and induction of lipogenic genes such as sterol regulatory element binding protein-1c (SREBP-1c) and fatty acid synthase in the liver of HFD-fed mice. We also found that CKS suppressed high glucose-induced lipid accumulation in HepG2 cells. CKS inhibited the high glucose-induced fatty acid synthase (FAS) and sterol regulatory element binding protein-1c (SREBP-1c) expression. Blood biochemical analyses and histopathologic examinations showed that CKS prevented liver injury. Moreover, the using pharmacological AMPK inhibitor revealed that AMPK is essential to suppression of SREBP-1c expression in CKS-treated cells. These results indicate that CKS prevents high glucose-induced lipogenesis in HepG2 cells by blocking the expression of SREBP-1c, FAS through AMPK activation, suggesting that CKS is a novel AMPK activator with a potential role in the prevention and treatment obesity.
1312 Platycodi radix Attenuates Dimethylnitrosamine-Induced Liver Fibrosis in Rats.

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Liver fibrosis is a wound healing response to a variety of chronic stimuli, including alcohol intake, viral infection, drugs, and metabolic disease. This study investigated the anti-fibrotic effects of the aqueous extract of the Platycodi Radix root (Changkil: CK) on dimethylnitrosamine (DMN)-induced liver fibrosis in rats by inducing Nrf2-mediated antioxidant enzymes. Repeated DMN exposure causes chronic liver injury with necrosis, fibrosis, and nodular regeneration through metabolic activation of CYP2E1 in experimental animals. CK inhibited DMN-induced increases in serum ALT and AST activities, fibrosis score, and hepatic malondialdehyde and collagen content. CK also inhibited DMN-induced reductions in rat body weight. CK inhibited DMN-induced increases in MMP-13, TIMP-1, and TNF-α mRNA and collagen type I and collagen smooth muscle actin protein. DMN-induced COX-2 expression and NF-κB activation was reduced by CK treatment. Furthermore, CK induced activation of Nrf2-mediated antioxidant enzymes such as γ-glutamylcysteine synthetase (γ-GCS), heme oxygenase-1 (HO-1), NAD(P)H quinone oxidoreductase 1 (NQO1), and glutathione-S-transferase (GST) in HepG2 cells. These results demonstrated that CK attenuates DMN-induced liver fibrosis through the activation of Nrf2-mediated antioxidant enzymes.

1313 Protective Effects of Saponins Isolated from the Root of Platycodon grandiflorum against Ovalbumin-Induced Airway Inflammation in a Mouse Allergic Asthma Model.

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Bronchial asthma is a chronic airway disorder characterized by airway inflammation, mucus hypersecretion, and airway hyperresponsiveness. This study investigated the protective effects of saponins isolated from the root of Platycodon grandiflorum (Changkil saponins: CKS) on airway inflammation induced by allergic reaction in mice. Mice were sensitized and challenged with ovalbumin (OVA) developed inflammation and remodeling in airway. CKS inhibited OVA-induced number of inflammatory cells and levels of TNF-α, IL-4, IL-5, IL-13, monocyte chemoattractant protein-1 (MCP-1) and OVA-specific IgE in bronchoalveolar lavage fluid. Also, CKS attenuated OVA-induced mucus hypersecretion in lung histopathological studies. CKS inhibited OVA-induced mRNA expression level of MMP-2, MMP-9, and MUC5AC in lung tissues. Furthermore, CKS blocked NF-κB p65 nuclear translocation in the nuclear extracts from lung tissues of OVA-challenged mice. These results suggest that CKS ameliorates OVA-induced inflammation and remodeling in airway.

1314 Topical Application of Cultivated Ginseng Suppresses 2, 4-Dinitrochlorobenzene-Induced Atopic Dermatitis in NC/Nga Mice.

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Atopic dermatitis (AD) is known to be related with local and systemic immunologic dysfunction that leads to Th1/Th2 response as well as being related to environmental factors. This study examined the inhibitory effects of 1-chloro-2,4-dinitrochlorobenzene (DNCB)-induced AD by cultivated ginseng (CG) (Panax ginseng C.A. Meyer) in NC/Nga mice. CG, an herb used in Korean herbal medicine, has been widely used in China and Japan to treat fatigue and to enhance resistance to many diseases. CG ameliorates DNCB-induced skin dermatitis severity, serum level of IgE and TARC, and mRNA expression of T helper (Th1) and Th2 cytokines in mice. In addition, CG reduced thickness of the dermis and dermal infiltration of inflammatory cells in histopathological examination. These results indicate that CG inhibits allergic contact dermatitis through the modulating of Th1 and Th2 responses and diminishing the inflammatory cells in the skin lesions in NC/Nga mice.

1315 Pleurotus eryngii Extracts Inhibits 2, 4-Dinitrochlorobenzene-Induced Atopic Dermatitis in NC/Nga Mice by the Regulation of Th1/Th2 Balance.

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Atopic dermatitis (AD) is a chronic, relapsing and inflammatory skin disease in humans and animals, caused by a complex interrelationship among genetic, environmental, pharmacologic, psychological, immunologic and skin barrier dysfunction factors. This study investigated the inhibitory effect of the Pleurotus eryngii extracts (PEE) on 2,4-dinitrochlorobenzene (DNCB)-induced atopic dermatitis (AD)-like skin lesions. Pleurotus eryngii has been used in traditional medicine for nutritional and medicinal food that enhances the host immune response as to various diseases. This study evaluated skin dermatitis severity, ear thickness, histopathological examination, and cytokines level in DNCB-applied mice treated with PEE. Continuous treatment of PEE inhibited the development of the AD-like skin lesions. PEE attenuates DNCB-induced skin dermatitis severity, serum level of IgE and TARC, and mRNA expression of TNF-α, INF-γ, IL-4, IL-5, and IL-13 in mice. Also, PEE reduced thickness of the dermis and dermal infiltration of inflammatory cells and mast cells in histopathological examination. These results suggest that PEE inhibits allergic contact dermatitis through the modulating of T helper (Th1) and Th2 responses and diminishing the inflammatory cells and mast cells infiltration in the skin lesions in NC/Nga mice.

1316 Chronic Oral Toxicity Study of the Extracts from Herbs in Phikud Navakot.

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Phikud Navakot is commonly used as Thai traditional medicine for alleviation hyperlipidemia, cardiovascular diseases, and cerebrovascular diseases. Chronic toxicity effects of the extracts from herbs in Phikud Navakot were performed because there were no 6-month toxicological studies reported in the literature. The repeated dose as 10, 100, 1,000 mg/kg/day of the extracts were randomly administered to both male and female Sprague Dawley rats as described in the OECD code 452 guideline. 183 rats survived to the end of the study while 20 rats died. The cause of death was considered to be related to the gavage technical error which positively correlated to high dose concentration. Mean body weights of dosed rats were similar to those of the control group. No differences in feed consumption, relative organ weights, and hematology and blood clinical chemistry were noted. Significant proportionally increased incidences of transitional cell hyperplasia of the renal pelvis in both sexes dosed rats. Therefore our obtained results suggested that Phikud Navakot is a relatively nontoxic herb for repeated oral administration. However the contraindication of the usage of Phikud Navakot is related to induce transitional cell hyperplasia of the renal pelvis after prolong high dose oral administration.

1317 Plum Consumption Modulate Gut Microbiota and Obesity in Zucker Rats.

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The role of peach and plum polyphenols on the modulation of gut microbiota and the possible relationship with obesity is unknown. We investigated how peach and plum consumption can modulate metabolic syndrome and relative proportions of gut microbiota populations in feces of obese Zucker rats. Experimental groups were fed a control or peach or plum juice ad libitum during 11 weeks, the control and lean groups received water with same amount of glucose than fruit juices. At the end of the study blood was analyzed for glucose, insulin triglycerides, cholesterol and LDL oxidation levels. DNA in feces was analyzed using 454-pyrosequencing and qPCR. Results showed that only plum juice consumption prevented weight gain and modulated microbiota populations in feces; this was related to the higher content of polyphenols in plum juice (3X of peach). However, both peach and plum exerted similar protective effects on metabolic syndrome and inflammatory markers.
Clarias gariepinus juveniles (mean weight (14.9±0.83 g) per tank, each in triplicate. Five treatment tanks were fed a diet containing 40% crude protein supplemented with varying inclusion of probiotic comprising about 109 colony-forming units per gram of diet (the probiotic diet). Diet T0 contain 0g probiotic (control diet) while the other group contain 0.5 g, 1.0 g, 1.5 g and 2.0 g probiotic diet. Results shows that Fish fed with diet T1 (0.5 g probiotic) had the best growth performance. There was no significant different (P<0.05) in the Mean Corpuscular Volume, Mean Corpuscular Haemoglobin and Mean Corpuscular Haemoglobin Concentration, of fish fed different concentration of probiotic. All blood parameter obtained were between the range of recommended fish blood. It is concluded that using probiotic (especially at 0.5 g) as supplementary feed on Clarias gariepinus showed a slight increase in the haematological parameters compare with the control diet but it has no negative impact on the health status of the species. However, probiotic (Lactobacillus and bifidobacterium) can be used as a probiotic agent in aquaculture, to enhance fish health, survival and growth performance.
1323 Relationship between Silver Nanoparticle Intracellular Accumulation and Cytotoxicity in L-929 Fibroblasts.

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Medical devices containing silver nanoparticles (AgNPs) may release NPs or leach silver (Ag) ions, both of which are cytotoxic in vitro at high concentrations. Given the complexities of NP in vitro dosimetry, proper assessment of AgNP cytotoxicity requires a better understanding of intracellular Ag concentration and observed bio-effects. Therefore, the objective of this study was to measure time-dependent AgNP cell internalization in L-929 fibroblast cells, a cell line that is commonly used for medical device biocompatibility testing, while simultaneously assessing cell viability. These endpoints were assessed following exposure of cells to 50 μg/ml AgNPs (10, 50, 100 and 200 nm) and to equimolar Ag ions. Intracellular uptake of AgNPs and Ag concentrations in L-929 cells were assessed using time-lapse confocal microscopy, flow cytometry and ICP-MS at 4, 16 and 24 hrs after treatment. Metabolic activity and plasma membrane integrity of the cells were assessed at 4, 16 and 24 hrs after treatment using the MTT assay and ethidium homodimer-1 dead cell indicator, respectively. A time-dependent, linear increase in intracellular accumulation of AgNPs and Ag concentrations were observed using confocal microscopy and ICP-MS. Flow cytometric evidence of AgNP uptake and accumulation in cells was also observed from changes in side scatter characteristics vs. forward scatter characteristics dot plots. The results of the MTT assay suggest that Ag ions and 10 nm NPs are highly toxic, compared to 50, 100 and 200 nm NPs. Membrane integrity analysis of time-lapse images revealed that 10 nm AgNPs are cytotoxic within 1 hr after dosing and all cells are dead after 10 hrs. In contrast, equimolar concentrations of Ag ions kill cells within an hour. This study demonstrated that the size-dependent in vitro cytotoxicity of AgNPs to L-929 fibroblasts correlated with intracellular Ag accumulation, and underscores the value of using multiple analytical methods when determining NP accumulation and cytotoxicity.

1324 Role of Sample Preparation in In Vitro Cytotoxicity Responses to Silver Nanoparticles.

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Metallic nanoparticles readily agglomerate in aqueous media and this effect can influence their physical properties and in vitro biological responses. The goal of this study was to compare effects of cell culture medium containing 10% FBS and DI water as suspension vehicles on AgNP size and agglomeration, and 2) compare effects of pre-mixing of medium and AgNPs on their biological responses in L-929 fibroblasts compared to direct addition of nanoparticles to the cells. To assess the effects of premixing on agglomeration, AgNPs were pre-mixed with cell culture medium or DI water for 1 min, or 1, 5, 24, or 120 hr. Results of dynamic light scattering analysis showed that pre-mixing AgNPs with medium maintained particle dispersion better than DI water. The hydrodynamic diameter of AgNPs increased proportionally to the pre-mixing time. AgNP agglomeration was size-dependent; 10 nm AgNPs agglomerated more readily than 100 nm and 200 nm particles. To assess the effects of premixing on biological responses, AgNPs pre-mixed with cell culture medium for 1 min, or 1, 5, or 24 hr were added to L-929 fibroblasts, or were added to the cells without premixing. After 24 hr exposure, cell viability was assessed by using the standard MTT assay. After 4 and 24 hr exposures, the degree of cell necrosis (via 7-AAD dye) and apoptosis (via Annexin V dye) was assessed by using the standard MTT assay. After 4 and 24 hr exposures, cell necrosis and apoptosis were assessed using 7-AAD and Annexin V dyes, respectively, with FC. Multiple FC controls (including cells alone, AgNPs alone, and single fluorescein controls and their combinations) were used to optimize the experiment and eliminate background autofluorescence and fluorochrome overlap. The data show that cell necrosis and apoptosis in AgNP-exposed fibroblasts depend on exposure time point and AgNPs size. Cells treated with 10 nm nanoparticles at 50 μg/ml showed 7- to 22-fold increases in percentage of apoptotic cells and 2- to 33-fold increases in percentage of necrotic cells after 4 hrs and 24 hrs, respectively. Cells treated with 200 nm nanoparticles at 50 μg/ml showed only up to 6 fold increases in degree of apoptosis after 24 hr. The data show that AgNPs produced a dose- and time-dependent decrease in cell viability; however, 200 nm AgNPs were significantly less toxic than smaller sized particles. Thus, standard FC assays can be utilized to assess apoptosis and necrosis in response to nanomaterial exposure.

1325 Flow Cytometry Evaluation of Cell Cytotoxicity Induced by Silver Nanoparticles.

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Particles possess unique properties in the nanoscale, e.g., enhanced catalytic activity, high surface area and surface energy, and light emission/absorption properties, which might result in interference with colorimetric in vitro cytotoxicity assays such as MTT, LDH release, and Neutral Red. Alternatively, assays that do not use spectrophotometric detection, such as trypan blue exclusion or flow cytometry (FC) based assays, are less likely to be influenced by nanoparticle interference. The aim of this study was to evaluate FC assays to assess the cytotoxicity of three different sizes (10, 100, or 200 nm) silver nanoparticles (AgNPs) at different mass concentrations (1, 25, or 50 μg/ml) in L-929 fibroblast cells. After 4 hrs and 24 hrs exposure, cell necrosis and apoptosis were assessed using Annexin V and 7-AAD dyes, respectively, with FC. Multiple FC controls (including cells alone, AgNPs alone, and single fluorescein controls and their combinations) were used to optimize the experiment and eliminate background autofluorescence and fluorochrome overlap. The data show that cell necrosis and apoptosis in AgNP-exposed fibroblasts depend on dose, exposure time, and AgNPs size. Cells treated with 10 nm particles at 50 μg/ml showed 7- to 22-fold increases in percentage of apoptotic cells and 2- to 33-fold increases in percentage of necrotic cells after 4 hrs and 24 hrs, respectively. Cells treated with 200 nm nanoparticles at 50 μg/ml showed only up to 6 fold increases in degree of apoptosis after 24 hr. The data show that AgNPs produced a dose- and time-dependent decrease in cell viability; however, 200 nm AgNPs were significantly less toxic than smaller sized particles. Thus, standard FC assays can be utilized to assess apoptosis and necrosis in response to nanomaterial exposure.

1326 Incorporation of Silver Nanoparticles into a Degradable Poly(L-Lactide-Co-Epsilon-Caprolactone) Copolymer Scaffold for Skin Regeneration.

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The development of an antibacterial, degradable scaffold system that utilizes patient-derived cells would improve upon current skin grafting techniques which often result in severe scarring, aesthetically undesirable mismatches in skin tones, and are susceptible to surgical site infection. The objective of this study was to characterize and assess the toxicity of an electrospun scaffold of poly(L-lactide-co-epsilon-caprolactone) (PLCL) incorporating antibacterial 20nm silver nanoparticles (AgNPs). The content and distribution of 20nm AgNP incorporated within the PLCL scaffold was optimized to maximize their biocompatibility and antimicrobial activity. The toxicity of the scaffold to human epidermal keratinocytes (HEK) was assessed using live/Dead and alamarBlue viability assays following 7 days and 14 days. No significant decreases in cell viability were noted at either time point and cell proliferation increased 120% at 7 days and 200% at 14 days on both control and AgNP incorporated scaffolds. After 14 days, scanning electron microscopy revealed a confluent layer of HEK on the surface of the scaffolds, and fluorescent microscopy confirmed cell migration into the scaffold interior. The antibacterial efficacy of the scaffold was evaluated against Escherichia coli and Staphylococcus aureus. The mechanical properties of the PLCL scaffold were assessed via uniaxial tensile testing to failure. A slight decrease in the modulus of elasticity was observed following AgNP incorporation compared to control, while cellular attachment increased the modulus of elasticity significantly. (Supported by NIH RO1 ES016138)

1327 Formation of a Protein Corona on Silver Nanoparticles Mediates Cellular Toxicity via Scavenger Receptors.

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Addition of a protein corona (PC) on the surface of nanomaterials can modify their activity, bio-distribution, cellular uptake, clearance and toxicity. As silver nanoparticles (AgNP) are incorporated into many products as anti-bacterial/fungal agents, the risk of human exposure escalates. We hypothesize that AgNPs will associate with proteins commonly found in human serum and cell culture media forming PCs which will impact cell activation and cytotoxicity. Furthermore, we believe that
activation of scavenger receptor B (SR-B) mediates this toxicity. Citrate- or PVP-coated AgNPs were incubated with human serum albumin (HSA), bovine serum albumin (BSA), high-density lipoprotein (HDL), or water (control). AgNPs associated with each protein (HSA, BSA, HDL) forming PCs by TEM, UV-vis spectroscopy, and altered Z-potential and hydrodynamic size. Rat aortic endothelial (RAEC) and rat lung epithelial (RLE) cells were exposed to increasing concentrations of AgNPs (0.625, 1.25, 25, or 50 μg/ml) with or without PC. For 3h or 6h. All PC-coated AgNPs demonstrated a dose-response relationship in cytotoxicity in both cell types. To determine the role of SR-B in the observed cytotoxicity, cells were exposed to AgNPs with or without PCs for 3h in the presence of a SR-B antagonist. Treatment with the SR-B antagonist inhibited cytotoxicity in RAEC but not RLE. Lastly, cell activation was assessed at 1h by measuring interleukin-6 (IL-6) mRNA expression. All PC-coated AgNPs induced IL-6 mRNA expression at 1h in both cell types whereas treatment with the SR-B antagonist was found to inhibit expression. Differences in the induction of IL-6 were found between PC-coated AgNPs based upon suspension (citrate or PVP). This study characterizes a PC on AgNPs using proteins found in human serum and cell culture media. The presence of these PCs influenced cytotoxicity and cell activation through SR-B leading to altered cell responses. This work was supported by the U19 ES019525 and R01 ES09311.

1328 Importance of p38MAPK and pmk-1 a Caenorhabditis elegans Homologue, in Silver Nanoparticles-Induced DNA Damage Response and Apoptosis.


Silver nanoparticles (AgNPs) have recently received much attention for their possible applications in new material design, biotechnology and other commercial purposes. Our previous studies conducted in Jurkat T cells and Caenorhabditis elegans displayed that AgNP exposure caused dose and time dependent increase in only p38MAPK and pmk-1 expression among all other stress responsive proteins and DNA damage in Jurkat T cells. These findings motivated to ask whether and how p38MAPK and pmk-1 is involved in AgNP induced DNA damage response and apoptosis. This work was supported by the U19 ES019525 and R01 ES09311.

1329 Cytotoxicity and Genotoxicity of Silver Nanoparticles and Silver Ions to CHO K1 Cells.

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Silver nanoparticles (Ag NPs) are used in a variety of commercial products due to their antimicrobial activity. As both in vitro and in vivo studies have demonstrated toxic effects of Ag NPs, there is an urgent need to explore the toxicity of Ag NPs. Here we studied the cytotoxicity and genotoxicity of Ag NPs (BSA coated, 15.9±7.6 nm) and silver ions (Ag+) to Chinese hamster ovary (CHO K1) cells. To analyze the cytotoxic effects the mitochondrial activity was determined by the MTT assay, intracellular reactive oxygen species (ROS) and the cell cycle were analyzed by flow cytometry. Fluorescence microscopy and flow cytometry based micronucleus assays were applied to quantify and quantify the number of micronuclei induced by Ag NPs and Ag+. P53 postlabeling was performed to detect DNA adducts. In addition, inductively-coupled plasma mass spectrometry (ICP-MS) and transmission electron microscopy (TEM) were applied to study the uptake and intracellular distribution of Ag NPs, respectively. A time and dose dependent decrease in mitochondrial activity and increase of intracellular ROS and Ag+ induced a cell cycle arrest in the G2/M phase. Micronucleus assay and P53 postlabeling revealed that both Ag NPs and Ag+ induced micronuclei and DNA adducts. Using TEM observations Ag NPs were found to be located in endosomes/lysosomes suggesting that Ag NPs are taken up by receptor mediated endocytosis. However, no Ag NPs were found in the nucleus suggesting that Ag NPs are presumably dissolved into Ag+ in the endosomes/lysosomes and released to the cytoplasm. From here they can enter mitochondria and/or the nucleus leading to an increased intracellular ROS level and the induction of DNA damage.

PS

1330 Chronic Exposure to Realistic Doses of Silver Nanoparticles Demonstrated Differential Cellular Responses Than Acute Exposure in Human Keratinocytes.

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One obstacle plaguing the field of nanotoxicology is the development of a mechanism to translate acute exposure data to an accurate prediction of real world implications of nanomaterials (NM). In an effort to enhance the efficacy of information gathered about Ag in vitro exposure to chronic NM exposure, a chronic NM dosing protocol was designed and implemented in which human keratinocyte cells (HaCaT) were dosed with 50 nm silver nanoparticles (Ag-NP) 8 h a day, 5 days a week, for 3 months. Working concentrations were based off the permissible exposure limits set by OSHA and were 0.4, 4, and 400 μg/ml. The HaCaT stress response of the chronically dosed cells was directly compared to a 24 h acute exposure at a concentration equal to the cumulative Ag-NP dosage encountered over the 3 months. Cellular endpoints evaluated included activate of Heat shock protein-27 signal transduction, ki67 expression, pro-inflammatory cytokine secretion, actin inflammation, and alterations in gene regulation. Results indicated that the chronically dosed HaCaT cells were functioning under sustained, augmented cellular stress; as seen with increased reactive oxygen species levels, HSP-27 signaling, cytokolic ki67 expression, and actin inflammation and disorganization. Furthermore, considerable IL-6 secretion was observed throughout the experiment, indicating a continuous inflammatory response. Most notably, these stress indicators were all significantly higher in chronically dosed cells vs. their acute counterparts, demonstrating a more severe response. Additionally, chronically dosed cells demonstrated a vastly higher modification to gene regulation, again representing the potential for a serious long-term impact. In conclusion, this study identified a significant variation in the HaCaT stress response following chronic exposure of Ag-NPs vs. an acute scenario and offers a novel approach to nanotoxicology research.

1331 Evaluation of Uptake and Cytotoxicity of Citrate Stabilised Gold Nanoparticles in Chinese Hamster Ovary (CHO) Cells prior to the In Vitro Mammalian Mutation Test.

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The health risk assessment of engineered gold nanoparticles (AuNPs) requires the generation of hazard identification data. Currently the Organisation for Economic Co-operation and Development (OECD) has released a guidance manual for the testing of manufactured nanomaterials. Within this document various established toxicological endpoints have been proposed for study, including the in vitro mammalian cell gene mutation test (OECD TG476) for genotoxicity assessment. Prior to this mutation study, preliminary experiments were required to assess the cytotoxicity and uptake of the AuNPs into the cells. Chinese Hamster Ovary (CHO) cells were treated with 14 nm, 20 nm, and 40 nm citrate stabilised gold nanoparticles. The Roche xCELLigence RTCA system was used to determine cytotoxicity of the particles into the cells. The nanoparticles were non-toxic for the duration of the period tested. The use of this technology eliminated any potential interference of the nanoparticles with the assay. The uptake of the particles into the cells was visualised using the CytoViva Hyperspectral Imaging system and showed a time-dependent uptake of the particles into the cells. This preliminary data confirms lack of toxicity of the particles prior to use in the mutation assay, and confirms uptake of the particles into the cells. Mutagenicity studies using this cell line will be conducted to measure mutation at hypoxanthine-guanine phosphoribosyl transferase (HPRT). Although gold nanoparticles have been shown in literature to induce some DNA damage; to our knowledge, the genotoxicity of gold nanoparticles has not been assessed using the HPRT mutation assay to date.
The majority of studies have focused on nanoparticle (NP) fate and their induced damage with limited information on the physiological environment’s role in altering NP properties. Artificial fluids (AFs) have been used to test the bioavailability of metallic compounds (difference between the amount of a substance a person is exposed to and the amount of substance the body receives). Literature has demonstrated that inhaled NPs accumulated in lungs and clearance was hindered during chronic exposure indicating the observed effects should become more pronounced over time due to impeded clearance. Based on these findings, this study examined the impact of artificial intestinal, alveolar, lysosomal, and gastric fluid on the physical properties of hydrocarbon coated silver (Ag-HC) and polycarbonate coated silver (Ag-PS) NPs and the associated changes in toxicity. Since inhalation exposure is of concern, the toxicity of the Ag-NPs exposed to the AFs was evaluated in alveolar macrophages and doses represent a week or year of exposure (0.5 ng/ml and 25 ng/ml) based on the concept that this delayed clearance will result in macrophages continually being recruited to engulf the NPs. For all the AFs, the Ag-HC NPs demonstrated large agglomeration, limited ionic dissociation, and no changes in cell viability. Ag-PS NPs exposed to alveolar and intestinal fluid demonstrated a similar trend, in addition to a loss of the PS coating. For the gastric fluid, large agglomerates were observed but these were rare and most likely due to the Ag NPs precipitating as AgCl. Interestingly, the Ag-PS NPs exposed to lysosomal fluid demonstrated a loss of coating, less agglomeration, and significant decreases in cell viability. Since NPs are most likely taken up via endocytosis in cell cultures or by phagocytic cells in vivo, the interactions of the NP with the lysosomal environment has critical implications on mediating the NP cellular consequences. Based on this study, the lysosomal environment has the potential to make a NP more toxic over time. (88ABW-2012-5185)

Silver nanoparticles (Ag-np) are distinctively reported to be toxic to mammalian cells. Less is known about the signalling response triggered in cells to counteract such toxicity. This study was initiated to enhance our mechanistic insight on correlation between, DNA-dependent protein kinase catalytic subunit (DNA-PKcs) and toxicity of Ag-np. The toxicity of polyvinyl alcohol (PVA)-coated Ag-np was studied in normal human lung fibroblast and human breast cancer and brain cancer cells. DNA-PKcs inhibition was carried out to investigate impact of DNA-PKcs on influencing toxicity of Ag-np. The toxicity was evaluated using changes in cell survival, DNA damage and repair and telomere length. We observed concurrent activation of DNA-PKcs and JNK pathway in cancer cells upon Ag-np treatment, which were anticipated as physiologic responses to DNA damage and repair. JNK pathway was insufficiently activated in DNA-PKcs inhibited cancer cells, abolishing the signalling events required for mediating DNA repair. Further investigation on genotoxic effect of Ag-np indicated that Ag-np causes telomere attrition and dysfunction in human cancer cells by disrupting shelterin complex integrity and telomerase expression. Recruitment of activated DNA-PKcs to damaged telomeres signifies the importance of DNA repair machinery at damaged telomeres. DNA-PKcs inhibition potentiates the damaging effect of Ag-np at telomeres in human cancer cells. Abrogation of JNK mediated DNA repair and substantial damages at telomeres lead to higher cell death in DNA-PKcs inhibited, Ag-np treated cancer cells. Altogether, presence of DNA-PKcs effectively reduces the toxic effects of Ag-np in human cancer cells by triggering the activation of JNK pathway and telomerase pathways.

The present study, the ion-release kinetics from citrate capped 20, 50, and 80 nm nanosilver dissolution in artificial physiological fluids: coating and pH impact on degree of ionic dissociation. L. Bravdich-Stolle, E. Breitner, E. I. Mauer, B. Stacy and S. M. Hussain, 711 HPW/RHD, Wright-Patterson AFB, Dayton, OH.

The majority of studies have focused on nanoparticle (NP) fate and their induced damage with limited information on the physiological environment’s role in altering NP properties. Artificial fluids (AFs) have been used to test the bioavailability of metallic compounds (difference between the amount of a substance a person is exposed to and the amount of substance the body receives). Literature has demonstrated that inhaled NPs accumulated in lungs and clearance was hindered during chronic exposure indicating the observed effects should become more pronounced over time due to impeded clearance. Based on these findings, this study examined the impact of artificial intestinal, alveolar, lysosomal, and gastric fluid on the physical properties of hydrocarbon coated silver (Ag-HC) and polycarbonate coated silver (Ag-PS) NPs and the associated changes in toxicity. Since inhalation exposure is of concern, the toxicity of the Ag-NPs exposed to the AFs was evaluated in alveolar macrophages and doses represent a week or year of exposure (0.5 ng/ml and 25 ng/ml) based on the concept that this delayed clearance will result in macrophages continually being recruited to engulf the NPs. For all the AFs, the Ag-HC NPs demonstrated large agglomeration, limited ionic dissociation, and no changes in cell viability. Ag-PS NPs exposed to alveolar and intestinal fluid demonstrated a similar trend, in addition to a loss of the PS coating. For the gastric fluid, large agglomerates were observed but these were rare and most likely due to the Ag NPs precipitating as AgCl. Interestingly, the Ag-PS NPs exposed to lysosomal fluid demonstrated a loss of coating, less agglomeration, and significant decreases in cell viability. Since NPs are most likely taken up via endocytosis in cell cultures or by phagocytic cells in vivo, the interactions of the NP with the lysosomal environment has critical implications on mediating the NP cellular consequences. Based on this study, the lysosomal environment has the potential to make a NP more toxic over time. (88ABW-2012-5185)
AgNPs in dilutions of an environmentally relevant freshwater (30 μS/cm and 150 μS/cm) were examined and related to the associated impact on an aquatic organism (Daphnia magna). Diluted suspensions of nanoparticles were placed on a multi-tube vortexer and orbital shaker for 0, 1, 2, 3 and 7 days. The acute toxicity of the AgNPs suspensions was then assessed with D. magna at 0 and 7 days post interaction between the particles and test media. An increase in hydrodynamic diameter measured by dynamic light scattering and field flow fractionation over time was observed at a relatively higher specific conductivity of 150 μS/cm in 20nm particles (3.3 fold increase) and only a small increase in 50 and 80nm particles (1.4 and 1.2 fold increase, respectively). At a lower conductivity of 30 μS/cm a 1.7, 1.0, and 1.2 fold increase was observed in 20, 50 and 80nm, respectively. Results showed that although the total concentration of silver in solution decreased with time, there was a consistent spike in dissolved concentration after 2-3 days interaction, followed by a steady decrease in dissolved silver in 150μS/cm and 30 μS/cm medium. This suggests that the concentration of dissolved silver in environmentally relevant ionic strength media increases over time after the introduction of capped AgNPs which may have implications on their antimicrobial properties. When exposed D.magna was exposed to 150μS/cm and 30μS/cm test media, 30μS/cm test media induced more toxicity than 150μS/cm test media. Toxicity increased with longer nAg interaction time with smaller particles inducing more toxicity than larger particles.

1337 Silver Nanoparticles Induce Mast Cell Degranulation via Scavenger Receptors.

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Silver nanoparticles (AgNPs) are increasingly incorporated into a variety of consumer and industrial products such as water filters and cosmetics for their antimicrobial properties. This has increased human exposures to AgNPs and therefore the possibility of adverse health effects. Mast cells are well known to orchestrate allergic immune responses through degranulation and release of pre-formed mediators such as histamine. Furthermore, mast cells have been shown to mediate pulmonary inflammation following exposure to nanoparticles in a murine model. We therefore examined whether AgNPs could induce mast cell degranulation. Bone marrow derived mast cells (BMMCs) were generated from femoral bone marrow of C57BL/6 mice. BMMCs were exposed to either citrate- or polyvinylpyrrolidone (PVP)coated AgNPs (20 nm or 110 nm diameter) at increasing concentrations (6.25, 12.5, 25, or 50 μg/ml) for 3, 6, or 24 h. Exposure to 20 nm AgNPs, but not 110 nm AgNPs, was found to cause concentration-dependent degranulation of BMMCs at 1 h. TNF-α gene expression was increased in BMMCs following AgNP exposure while TNF-α protein levels were increased only after exposure to citrate-coated AgNPs at 50 μg/ml. To determine the mechanism of BMMC degranulation following exposure to 20 nm AgNPs, we examined scavenger receptors which have been shown to mediate nanoparticle uptake in macrophages. PCR and flow cytometry demonstrated the presence of scavenger receptor class B1 (SR-B1) on the surface of cultured BMMCs. To determine the role of SR-B1, BMMCs were treated with two different SR-B1 antagonists (blt-1 or blt-2). Treatment with either SR-B1 antagonist was found to prevent AgNP-induced degranulation of BMMCs. These in vitro findings suggest that AgNPs may induce an inflammatory response via mast cell degranulation in vivo, which is dependent upon nanoparticle size and scavenger receptor activation. Therefore mast cell degranulation may be considered as a indicator of nanomaterial toxicity. This work was funded by NIEHS R01 ES019311 and U19 ES019525.

1338 High-Throughput Methods for Assessing the Molecular Toxicity of Nanomaterials in Bacteria.

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Synthesis and use of nanoparticles has skyrocketed during the past decade. To ensure that nanotechnology is safely and sustainably developed, rapid, cost-effective methods are needed to determine the toxicity of nanoparticles. Here, we report the application of a suite of sub-lethal assays as well as a growth inhibition assay to a series of silver and metal oxide nanoparticles in bacteria (Escherichia coli). Sub-lethal effects such as perturbation of membrane integrity (using PI/STYO) disruption of membrane potential (using DBAC and reduction of respiration rate (using XTT) were used to identify toxic nanoparticles. To further look into mechanism of toxicity, intracellular Reactive Oxygen Species (ROS) and the intrinsic property of nanoparticles to oxidize electron (abiotic ROS generation) were measure by using H2DCFDA and DCFH-DA was examined and related to the associated impact on an aquatic organism (Daphnia magna). Diluted suspensions of nanoparticles were placed on a multi-tube vortexer and orbital shaker for 0, 1, 2, 3 and 7 days. The acute toxicity of the AgNPs suspensions was then assessed with D. magna at 0 and 7 days post interaction between the particles and test media. An increase in hydrodynamic diameter measured by dynamic light scattering and field flow fractionation over time was observed at a relatively higher specific conductivity of 150 μS/cm in 20nm particles (3.3 fold increase) and only a small increase in 50 and 80nm particles (1.4 and 1.2 fold increase, respectively). At a lower conductivity of 30 μS/cm a 1.7, 1.0, and 1.2 fold increase was observed in 20, 50 and 80nm, respectively. Results showed that although the total concentration of silver in solution decreased with time, there was a consistent spike in dissolved concentration after 2-3 days interaction, followed by a steady decrease in dissolved silver in 150μS/cm and 30 μS/cm medium. This suggests that the concentration of dissolved silver in environmentally relevant ionic strength media increases over time after the introduction of capped AgNPs which may have implications on their antimicrobial properties. When exposed D.magna was exposed to 150μS/cm and 30μS/cm test media, 30μS/cm test media induced more toxicity than 150μS/cm test media. Toxicity increased with longer nAg interaction time with smaller particles inducing more toxicity than larger particles.

1339 Role of microRNA in Toxicity of Silver Nanoparticles in Jurkat T Cells.


In this study, we intend to identify whether miRNAs are involved in toxicity of silver nanoparticles (AgNPs), we investigated miRNA expression in AgNPs treated Jurkat T cells using miRNA and miRNA microarrays. Surprisingly small numbers of genes were affected by both silver exposures(19 up- and 2 down-regulated miRNA by AgNPs exposure, and 2 up- and 1 down-regulated mRNA by Ag ion exposure). More important numbers of differentially expressed (DE) miRNAs than DE mRNAs were revealed by miRNA microarray, such as, 31 up- and 51 down-regulated miRNAs by AgNPs exposure, whereas 11 up- and 27 down-regulated miRNAs by Ag ion exposure. The most dramatic alteration was observed in has-miR-1238 (a decrease of 67 folds) by AgNPs exposure. An integrated analysis of miRNA and mRNA expression was conducted on DE miRNA and DE mRNA by AgNPs and Ag ion exposure, which revealed that the expression of miRNA, hsa-miR-219-5p, was negatively correlated with that of mRNA, metallothionein 1F (MT1F) and tribbles homolog 3 (TRIB3), in the cells exposed to AgNPs; whereas, the expression of has-miR-654-3p was negatively correlated with the expression of mRNA, endonuclease G-like 1 (EDGGL1) in the cells exposed to Ag ions. However no miRNA regulated by has-miR-1238 was identified. Individual validation was followed on these 3 miRNA-mRNA pairs using qPCR. Bioinformatics analysis was further conducted using Pathway Studio, which suggests hsa-miR-219-5p - MT1F and -TRIB3 pairs play an important role in AgNPs toxicity in Jurkat T cells by regulating the genes in important stress response pathways, such as, cell survival, cell death and oxidative stress. Overall results suggest that compared to serious toxicity, only limited genes were affected by AgNPs at transcriptional level, whereas more important number of miRNAs were identified, which suggests fine regulation with miRNA may play important role in AgNPs toxicity.

1340 Assessing the Influence of Gold and Silver Nanoparticles on Conventional In Vitro Cytotoxicity Assays.

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Gold nanoparticles (AuNPs) and silver nanoparticles (AgNPs), due to their optical properties are used in optics and biomedical applications. A principal property of these NPs is their enhanced surface plasmon resonance which enables them to scatter and absorb light with great efficiency. When assessing NP toxicity, there are many challenges as conventional methods are optimally biased and make use of absorbance, luminescence and fluorescence. The objective of this work was to assess if AuNPs and AgNPs are capable of interfering with currently accepted in vitro cyotoxicity assays. Changes in the endpoints of the XTT, ATP, LDH and MitoPT JC-1 assays was assessed in the absence and presence of cells, after the addition of AuNPs and AgNPs, to assess interference. Results obtained from XTT, ATP and LDH assays in the presence of cells showed toxicity. In the absence of cells, the XTT and ATP assays showed significant changes in their absorbance and luminescence; this could be explained by product adsorption to the NP or interference of NPs with the readings of the products assessed. On the other hand, the LDH assay showed little or no change in fluorescence. When using the MitoPT JC-1, evaluation of cell treated with the NPs showed viable mitochondria, however in the absence of cells there is a decrease in the fluorescence which too can be explained by product adsorption to the NP. In conclusion, NPs have the ability to interfere in vitro toxicity assays, skewing results. For this reason when assessing NP toxicity using these assay systems, their reliability and sensitivity needs to be determined. To overcome this interference of the gold and silver NPs a non invasive system such as the xCELLigence RTCA system which does not use absorbance, luminescence, and fluorescence, can be used to determine the viability of cells by measuring the cell index, a dimensionless parameter, which is an expression of cell adherence.
Traditional in vitro toxicology methods require dispersion of nanomaterials (NMs) in biological media for administration to cells, which does not depict realistic inhalation exposure. The objective of this work is to design and optimize a system to mimic inhalation exposure by delivering well-characterized NM aerosols to cells grown at the air-liquid interface. Gold or silver NMs were drawn into the gas phase from an aqueous dispersion using electrospray. The aerolized NMs were introduced into a chamber and deposited onto cells using electrostatic deposition. The deposition of NMs in the chamber was assessed by evenly placing copper grids throughout the chamber and imaging particle deposition using transmission electron microscopy (TEM). NM deposition on cells grown in the chamber was quantified by digesting the samples and measuring gold or silver content using inductively coupled plasma – mass spectrometry (ICP-MS). A human nasal epithelial cell-line was grown in the chamber, and the viability was assessed using the Alamar Blue assay. Results showed that gold and silver NMs could be deposited uniformly in the chamber and that the dose could be controlled by varying the electric field strength and frequency used for electrostatic deposition. Additionally, the dose was found to be relevant compared to NM deposition in the respiratory tract predicted by the Multiple Path Particle Dosimetry model. The results of the Alamar Blue assay showed that the cells could be sustained in the chamber, and the application of electric fields did not have an effect on cell viability. This study demonstrates a promising step forward in the development of a standardized realistic exposure method for assessing NM toxicity in vitro.

Nanoparticle size (nAg) is the most widely used engineered nanoparticle in consumer products, so much attention is being given to environment, health, and safety (EHS) risk associated with nAg exposure. The purpose of this study was to evaluate the agglomeration and dissolution of engineered nAg in artificial biofluids to better understand nAg stability throughout the body. Twenty-four hour time-course evaluations of 1 mg/L nAg in artificial biological fluids (alveolar, intestinal, gastric [pH 1.5], and intestinal fluid) were conducted using field flow fractionation-inductively coupled plasma-mass spectrometry (FFF-ICP-MS), dynamic light scattering (DLS), spectrophotometry, and dissolution methods. Silver nitrate (AgNO3) was used as a positive control. Citrate- and PVP-Ag (20 nm) in alveolar fluid and intestinal fluid showed similar settling patterns, with absorbance levels decreasing only 10% from t=0 to t=24h, likely due to the presence of surfactants. In contrast, these same particles in interstitial fluid and gastric fluid (pH 1.5) show a 40% decrease in absorbance at 24 h. Rapid agglomeration of coated nAg appeared to be occur, as indicated by the sudden formation of black particulates upon addition of the coated-nAg to all artificial biofluids tested, nAg showed marked size decrease over 24 h in interstitial, intestinal, and gastric fluids. The decrease in surface area resulting from this agglomeration may also have an effect on Ag dissolution rates. Citrate- and PVP-coated nAg dissolution, as well as AgNO3, suggests the loss of Ag+ when compared with Ag+ concentrations in water. The high concentration of chloride ions in many artificial biofluids may lead to the rapid formation of AgCl. These results demonstrate that nAg is likely to undergo several dissolution and agglomeration changes as the particle moves through different organ systems. Furthermore, these data can be used to generate prediction models of nAg dispersion throughout the body.

Manufactured silver nanomaterials (AgNPs) are used as antimicrobials in many consumer products. Although increased use of AgNPs increases risk of exposure by inhalation or ingestion, there are few data on human health risks associated with exposure. Here, we evaluated the toxicity of AgNPs in the murine macrophage J774A.1 cell line. Macrophages play a key role in the inflammatory response by phagocytosis of pathogens, debris, and particles. Phagocytosis of AgNPs by macrophages could expose cells to Ag and alter cell structure and function. We used two in vitro cytotoxicity assays, lactate dehydrogenase (LDH) and reduction of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), to compare cytotoxic effects of ionic Ag, polyvinylpyrrolidone PVP coated-AgNPs (MHD=13 nm), hydrogen reduced H2-AgNPs (14 nm), and citrate-AgNPs (12 nm). Cells were exposed to Ag, PVP-coated AgNPs, H2-AgNPs, or citrate-AgNPs, either in media alone or media supplemented with 1% fetal bovine serum (FBS) for 1, 4, or 24 hours before assessment. Each AgNP diminished MTT reduction capacity of J774A.1 cells with 50% reductions in activity in the low parts per million of Ag concentration range. Compared with AgNPs, Ag was a more potent cytotoxicant. LDH leakage increased after exposure to Ag and AgNPs indicated that all compounds produced damage to cell membranes. In MTT assays, addition of 1% FBS to media mitigated cytotoxic effects of all forms of Ag. In LDH assays, addition of FBS did not affect Ag-dependent membrane damage. These results indicate AgNPs affect integrity of cell membranes and metabolic competency of cells, although it is yet unclear whether these effects are mediated by phagocytosis of AgNPs or by accumulation of Ag solubilized from nanoparticles. Future studies that examine the disposition, fate, and effects of AgNPs will provide more information for assessment of bioavailability and potential human health risks. (This abstract does not reflect U.S. EPA policy.)

Nano-sized silver (nAg) is the most widely used engineered nanoparticle in consumer products, so much attention is being given to environment, health, and safety (EHS) risk associated with nAg exposure. The purpose of this study was to evaluate the agglomeration and dissolution of engineered nAg in artificial biofluids to better understand nAg stability throughout the body. Twenty-four hour time-course evaluations of 1 mg/L nAg in artificial biological fluids (alveolar, intestinal, gastric [pH 1.5], and intestinal fluid) were conducted using field flow fractionation-inductively coupled plasma-mass spectrometry (FFF-ICP-MS), dynamic light scattering (DLS), spectrophotometry, and dissolution methods. Silver nitrate (AgNO3) was used as a positive control. Citrate- and PVP-Ag (20 nm) in alveolar fluid and intestinal fluid showed similar settling patterns, with absorbance levels decreasing only 10% from t=0 to t=24h, likely due to the presence of surfactants. In contrast, these same particles in interstitial fluid and gastric fluid (pH 1.5) show a 40% decrease in absorbance at 24 h. Rapid agglomeration of coated nAg appeared to be occur, as indicated by the sudden formation of black particulates upon addition of the coated-nAg to all artificial biofluids tested, nAg showed marked size decrease over 24 h in interstitial, intestinal, and gastric fluids. The decrease in surface area resulting from this agglomeration may also have an effect on Ag dissolution rates. Citrate- and PVP-coated nAg dissolution, as well as AgNO3, suggests the loss of Ag+ when compared with Ag+ concentrations in water. The high concentration of chloride ions in many artificial biofluids may lead to the rapid formation of AgCl. These results demonstrate that nAg is likely to undergo several dissolution and agglomeration changes as the particle moves through different organ systems. Furthermore, these data can be used to generate prediction models of nAg dispersion throughout the body.

Particle size is thought to be a critical factor affecting the bioavailability of nanoparticles (NPs) following oral exposure. Nearly all studies of NP bioavailability focus on characterization of the primary particle size of the material as supplied or as dosed, and not on the effective particle size within the gastrointestinal tract (GIT), which is presumably most relevant for absorption. In the study reported here, agglomerative behavior of gold nanoparticles was evaluated throughout the gastrointestinal tract (GIT) using transmission electron microscopy (TEM). Agglomeration state within the GIT was then correlated with bioavailability as indicated by tissue levels of Au detected using inductively coupled plasma mass spectrometry (ICP-MS). Mice were dosed (10mg/kg) with either 22nm PEG-coated or uncapped Au NPs. Previous work done by this group using dynamic light scattering in simulated gastric fluid (SGF) has shown that uncapped Au NPs quickly agglomerate in this environment while PEG-coated Au NPs remain well dispersed after 24 hours in SGE. Samples were taken from various regions of the GIT at varying times.
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In recent years, the usage of nanoparticles has grown tremendously, and their potential to exert toxicity in aquatic environments is an increasing concern. This study evaluated the toxicity of 20 and 100 nm nanosilver and nanogold to Ceriodaphnia dubia in soft reconstituted water. The selected nanoparticles were chosen for nanosilver ability to dissolve in environmentally relevant solution and nanogold for its slow dissolution in environmentally relevant solution. Standard toxicity test have shown that Ceriodaphnia dubia are less sensitive to silver chronically than they are acutely. This is most likely a result of the addition of food binding to the silver reducing bioavailability. An altered testing procedure adding food after 8 hours from water renewal alongside standard testing procedures was evaluated acutely and chronically. This window of time will allow for the uptake of nanoparticles by the organisms that would otherwise not be available. In acute and chronic test Ceriodaphnia dubia were less sensitive to nanogold than nanosilver. When comparing acute and chronic 20 nm nanogold tests, 100% mortality was shown in the highest concentration 75 mg/L chronically with 60% survival acutely in the highest concentration. This would suggest the possibility that toxicity is occurring by different mechanisms. In 20 nm nanosilver chronic testing the altered testing procedure showed increase in mortality and delayed reproduction when compared to standard testing procedures confirming that the addition of food decreases sensitivity to the organism. Future work will include the testing of 100 nm nanosilver and nanogold. Further studies are needed to indicate that toxicity is occurring by different mechanisms when comparing acute-to-chronic ratios and testing of different aquatic organisms.
BAL cell oxidant production were measured in the Ag High group when compared to the Ag Low and DM groups. Following infection, LM lung burden increased significantly in the DM and Ag Low groups as compared to the Ag High group, peaking on day 6, with the highest burden in the DM group. LALN lymphocytes and BAL neutrophils, lymphocytes, and cellular oxidant production were elevated in the Ag High group on days 4 and 6 compared to DM and Ag Low groups. By day 8, LM lung burden, and BAL and LALN cell counts were similar for all groups. Induction of an early inflammatory response and oxidant burst in conjunction with increased lymphocyte proliferation in the lungs of the high dose AgNP group prior to infection enhanced the innate immune response and led to an increased clearance rate of bacteria from the lungs.

1351 Exposure to Inhaled Silver-Nanoparticles Results in the Induction of Gene Expression Alterations in Oxidative Stress Pathway Components, in a Time-Dependent Manner.


The use of silver nanoparticles (AgNPs) is ubiquitous, and they are now commonly employed in pharmaceuticals, medical imaging, medical devices, cosmetics, clothing, and other consumer products. Due to the commonality of AgNPs, along with a general lack of toxicity and safety testing, AgNPs have been deemed safe by de- fault. However, legitimate concerns exist, and a number of recent studies suggest that AgNPs pose a threat to human health. This study was conducted in order to examine the hypothesis that inhaled nano-sized Ag particles cause lung and extrapulmonary organ dysfunction via oxidative stress pathways. Male and female FVB/NJ mice were exposed to either 0.5 mg/m3 or 1 mg/m3 of AgNPs generated from metal rods, using a Pasal spark generator, via nose only inhalation for 4 hrs.

An n=5 animals were used in each treatment group, and all treated animals were compared to air exposed controls. At 2h, 24hr, 48hr, 72hr, and 72d-post exposure, animals were euthanized and blood, lavage and tissue were collected. RNA was isolated from lung and liver tissue and mRNA expression levels, of key oxidative stress genes, were quantified by real-time RT-PCR. A number of significant alterations in gene expression of key oxidative stress genes were observed in response to AgNP exposure. At 2 hours post exposure, Hmox1 and Keap1 expression increased ~2.75 fold in lung tissue, and modestly decreased to ~2.5 fold at 24 hours post exposure. Whereas in liver tissue, Keap1 expression changed from a 2.5 fold increase to a 3.5 fold increase in liver tissue, at 2 and 24 hours post exposure, respectively. Similarly, in liver tissue, Tmnd1 expression increased from 0.5 fold to a 2.5 fold increase at 2 and 24 hours post exposure, respectively. These experiments have suggested that exposure to inhaled AgNPs results in the induction of gene expression alterations in oxidative stress pathway components. Further, these alterations occur in a time-dependent manner.

1352 Genetic Influence of Pulmonary Response to Silver Nanoparticles Exposure.

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Within the last decade, nanotechnology has seen a boost in commercial, medical and industrial applications. However, there is great uncertainty in terms of the biological activity nanoparticles (NPs) may have on biological systems. In particular, this study sought out to determine the genetic influence on the pulmonary response to silver NPs. Murine models were employed to evaluate the variation of inter-strain response. Differences were quantitated by measuring inter-strain variability in terms of polymorphonuclear neutrophil (PMN) infiltration in bronchoalveolar lavage fluid. Initially, three mouse strains (C57BL/6J, FVB/NJ, and BALB/c) representing resistant, moderate, and sensitive responders, respectively, were exposed to 0.5 μg/g body weight of 20 and 110 nm citrate or PVP stabilized silver NPs by oropharyngeal aspiration. Bronchoalveolar lavage was performed 24 hr post exposure to measure PMN number and protein concentration. All three strains were found to be more sensitive to 20 nm silver citrate (40-85% PMN response) and 20 nm silver PVP (15-85% PMN response). We then performed a similar exposure in 8 strains of mice (DBA/2J, C57BL/6J, AKR/J, 129S1/SvJ, AJ/J, FVB/NJ, C3H/HeJ, Balb/cJ) at a lower dose of 0.25 μg/g body weight of 20 nm silver citrate. PMN responses between 1.5% and 60% were observed. In addition, temporal pulmonary response was measured in C57BL/6J, FVB/NJ, and BALB/cJ mice at 24 hrs and 7 days. At 24 hours, PMNs were found to be strain dependent, between 14-46%, and the response at 7 days was found to be between 0-15% PMN response. Based upon these initial findings on inter-strain response variation, it can be concluded that these mouse models show a susceptibility pattern among strains, and can help to elucidate sensitivity mechanisms in terms of exposure assessment and innate inter-species responses.
various cell-lines. However, the data on the toxicity of Ag-NPs in vivo is largely lacking. The main goal of this study was to determine the effect of two different diameter sizes (6nm, 10nm) silver nanoparticles on certain liver enzymes (alanine aminotransferase, aspartate aminotransferase and alkaline phosphatase) in serum and analysis of liver damage in Sprague-Dawley rats. Four groups of five male rats, each weighing approximately 80 ± 2 g, were administered orally, once a day for five days with doses of 5, 25, 50, 100 mg/kg body weight (BW) of Ag-NPs. A control group was also made of five rats. Serum was collected following standard protocols, and the activity of the liver enzymes (ALT, AST and ALP) was determined by colorimetric method. The results demonstrated that Ag-NPs exposure increased the activities of the liver enzymes (ALT, AST, ALP) and damage in liver tissue in exposed groups compared to control. The increase in the activity was larger in 6nm size AgNPs compared to 10nm size. Only the highest two concentrations 50 mg/kg and 100 mg/kg showed statistically significant increases in ALT and ALP in both diameter size AgNPs compared to control. AST activity showed an increase; however, it was not statistically significant compared to control. Furthermore, the smallest sized AgNPs (6nm size) had a greater ability to induce hepatic damage in Sprague-Dawley rats than the other sized AgNPs (10 nm). These data suggest that the AgNPs-induced hepatic toxicity effects against tissue cells are particle size-dependent, and thus, the particle size needs careful consideration in the design of the nanoparticle for biomedical uses.

1356 Increased Mucus Production and Histopathological Gill Alterations after Exposure to Nanosilver and Silver Nitrate.

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Silver nanoparticles are among the most widely used nanomaterials because of their antibacterial and antifungal properties. Despite their extensive use, information is now becoming available on the toxicity and fate of silver nanoparticles within living organisms. Mucus has both increased and decreased the toxicity of different xenobiotics by either concentrating the xenobiotic on the gills and body or encapsulating toxicants to prevent exposure. In order to understand the relationship between mucus, toxicity and silver exposure, zebrafish (ZF) and fathead minnows (FHM) were exposed to 36 or 96 hr to nominal 20 nanomolar AgNP- or citrate-coated silver nanoparticles (PVP-AgNPs; citrate-AgNPs) or silver nitrate (AgNO3) at 2 nominal concentrations (20 and 200 μg/L) or (2 and 6 μg/L), respectively. After 4 hr, ZF produced significantly more mucus secretion in every treatment than the control fish in a dose-dependent manner as measured by a phenol-sulfuric acid method. FHM gill RNA was extracted for microarray analyses. To quantify distribution of silver, skin, liver and GI were digested for ICP-MS. FHM gills were also paraffin embedded, sectioned and examined for histopathological alterations to pheno- typically anchor the molecular and chemical endpoints. The highest AgNO3 and both citrate-AgNPs concentrations caused significant atrophy in gill mucus goblet cells. Every silver treatment also had a significantly higher histopathological alteration index compared to control. Citrate-AgNPs (20 and 200 μg/L) had the highest incidence of alterations (3.5 and 3.25 times higher than control, respectively). The results suggest that mucus production and mucus-related cells are impacted by the particle size of AgNP and how the consumption of AgNP might disrupt the normal balance of microorganisms in the healthy G.I. tract. Supported by interagency agreements 224-12-0003 and AES12013 between the NCTR/FDA and NIEHS/NTP.

1357 Impact of Silver Nanoparticles (AgNP) on Bacteria Species Isolated from Gastrointestinal (G.I.) Tract of Sprague-Dawley (SD) Rats.

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The food industry is incorporating AgNP into a growing number of products, which suggests that consumers are being exposed orally to increasing amounts of these nanoparticles. The absorption of AgNP from the G.I. tract has been demonstrated; however the potential effects from the interactions of AgNP with bacteria of the G.I. tract are largely unknown. We measured the bacterial effects of AgNP on Lactobacillus sp. (G+) and Bacteroides sp. (G-) isolated from the G.I. tract of SD rats. Three sizes of AgNP (10, 75, and 110 nm) at concentrations of 0.1, 0.5, 1, 2, 5, 10 μg/ml were incubated with cultures of bacteria (in triplicate), and the bacterial effects at 60, 120, and 180 min were assessed using colony forming unit (CFU) and adenosine triphosphate (ATP) release assays. There were no CFUs in cultures of Lactobacillus sp. or Bacteroides sp. when incubated with any concentration of AgNP at concentrations of 1 μg/ml. There was a significant decrease in ATP at 10 and 110 μg/ml AgNP at concentrations of ≥2 μg/ml, suggesting enhanced bacterial effects by 10 nm AgNP. Both bacterial species showed a significant decrease in ATP released at 110 μg/ml (10, 75, or 110 μg/ml) at concentrations ≥1 μg/ml. At the equivalent number of cells (10^7 cells/ml), Bacteroides sp. were more sensitive than Lactobacillus sp. to the bactericidal properties of AgNP (10, 75, or 110 nm), likely reflecting differences in cell wall composition between species. Scanning electron microscopy showed that AgNP were attached to bacterial cells at the surface, suggesting that protein binding or cell wall distortion by AgNP might result in cellular toxicity or death. These results provide new insights into the bactericidal properties of AgNP and how the consumption of AgNP might disrupt the normal balance of microorganisms in the healthy G.I. tract. Supported by NIEHS P3 ES000210, ES016856 and Air Force Research Laboratory FA8650-05-1-5041.

1358 Influence of Dose, Size and Chemical Composition on Persistence of Silver Nanoparticles in the Rat Lung.


Silver nanoparticles (AgNPs) have antimicrobial activity and unique electrical properties, resulting in increasing use. While there have been studies of the biological effects of AgNPs, including persistence and clearance of AgNPs from the lung, an examination of the effect of AgNP size and surface coating (used to stabilize these materials when in solution) has not been fully investigated. We investigated lung deposition, retention and clearance of 20 nm and 110nm AgNPs coated with citrate or polyvinylpyrrolidone (PVP). Rats were instilled intratracheally with 0.1, 0.5 or 1.0 mg/kg of either one of the AgNP solutions or a vehicle control and were assessed at 1, 7 and 21 days post treatment. Ag was quantified in tissue using inductively coupled plasma mass spectrometry. At one day post instillation, lungs dosed with 1.0 mg/kg AgNPs had 0.25 to 0.51 mg Ag/g dry tissue and these amounts did not differ by surface coating or particle size. Sham controls had no detectable Ag. Total leukocytes and neutrophils in bronchoalveolar lavage fluid increased in a dose dependent manner 1-day after exposure to all AgNPs with significant increases in the 1.0mg/kg dose compared to all other doses for all 4 particle types. This increase persisted at the 7-day time point in all AgNP groups except for 20nm PVP. Distribution of Ag in the lung was determined using a multilabeled and semiquantitative scoring on paraffin-embedded lung sections to demonstrate Ag was preferentially localized to the bronchoalveolar duct junction at the 1-day time point. At 7-days post instillation, Ag was localized to the epithelial extracellular matrix of the terminal bronchioles. Uptake of Ag by alveolar macrophages was also observed. These findings suggest Ag can persist in the lungs over time and alveolar macrophages have a role in the clearance. Supported by U01ES02027 and P42ES00469. Nanomaterials supplied by the NIH NCNHR consortium.

1359 Subtle Surface Variations Influence Biological Compatibility of Gold Nanoparticles.

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To better understand the nanoparticle physicochemical properties that influence biocompatibility, we investigated how subtle surface variation on the surface of gold nanoparticles (AuNPs) influences biological responses. The zebrafish embryos were exposed to four distinct, well-defined ligand-stabilized 1.5 nm spherical AuNPs. The ligand shells for each AuNP contain: (1) anionic 2-mercaptoethanesulfonic acid (MES), (2) only cationic N,N,N-trimethylammoniummethanethiol (TMAT), (3) a mixed ligand shell containing MES and a small amount of triphenylphosphine (TPP), and (4) a mixed ligand shell containing TMAT and a small amount of TPP. Small-angle X-ray scattering (SAXS) confirmed the core size of the particles and that there was no agglomeration in the embryonic test medium. The only difference between partial and full functionalization was the P/Au ratio as determined by XPS analysis. The MES/TPP coated particles caused higher mortality and malformation than those fully functionalized with MES, and in TMAT/TPP coated particles a higher incidence of smaller size and pale gray eyes was observed. In the gene expression profiles of transcription factors for apoptosis (p53 and bax), eye development (pxa6a, otx2 and rxl) and pigmentation (sox10 and mir1), the gene expression of embryos were significantly disrupted in both mixed ligand particles in a dose-dependent manner, consistently with the observed developmental toxicity. In addition, laser ablation ICP-MS analysis revealed that biological absorption and distribution of AuNPs in exposed embryos were closely related to the toxicity phenotype and gene expression caused by differential surface functionalization. These data provide new insights into the understanding of how subtle surface changes impact biological compatibility and further emphasize the importance of material characterization. This research is supported by the NIH NCNHR consortium.
Gold nanoparticles (GNPs) possess unique physicochemical properties that may facilitate entry into the central nervous system (CNS) where they may act therapeutically. There is little information on GNPs biodistribution in specific brain regions or extent of inflammation induction. Experiments determined the localization and neuroinflammatory response of spherical GNPs (10 nm) after IV injection in male C57Bl mice. As a supplement, a known inflammogen, lipopolysaccharide (LPS, 2 mg/kg, sc), was tested. To determine the optimal buffer concentration to maintain GNP solubility, we measured aggregation of GNPs using various PBS concentrations (10, 1, 0.1, 0.01 X). 0.01X PBS produced the least amount of GNP aggregation and was used in all studies. The next experiment verified entry of GNPs into CNS. Mice were IV injected using the tail vein (200 μg/ml 10 nm GNPs in 0.01X PBS). After 24 hrs mice were perfused transcardially with 2% glutaraldehyde/paraformaldehyde and brains were collected. GNPs were measured using inductively coupled plasma mass spectrometry in whole brain homogenates. To specifically localize accumulation of GNPs in brain, septum, caudate, hippocampus, hypothalamus, cortex, frontal cortex, and spinal cord were dissected. Hypothalamus, hippocampus, and septum had the highest levels of GNPs (6.7, 6.2, and 4.6 μg Au/g, respectively). To evaluate brain inflammation, we used q-PCR analysis of frozen brain regions for study of pro-inflammatory mediators, IL-6, CCL2 and IL-1β. GNPs did not affect cytokine/chemokine expression in cortex, frontal cortex or hippocampus. LPS, as expected, caused a marked (100-fold) increase in the same cytokines. Results show that GNPs enter brain and concentrate in specific regions without eliciting an inflammatory response. Data raise the possibility of usefulness of GNPs in drug delivery and therapeutic treatment of CNS diseases.

The role of macrophage scavenger receptor A (SR-A) in the disposition of silver nanoparticles was investigated using cellular and animal models. Bone marrow derived macrophages from wild-type C57Bl/6J and SR-A deficient mice were exposed to silver nanoparticles (20 and 130 nm, primary size) coated with citrate and polyvinylpyrrolidone (PVP). Macrophages from wild-type mice demonstrated higher rates of silver binding/uptake compared those from SR-A(-/-) mice (0.5 vs. 0.003 hr⁻¹, respectively), which led to increased levels of cell associated silver (+10 fold at 24 hr) after exposure to 20 nm silver nanoparticles measured by inductively coupled plasma mass spectrometry (ICP-MS). In contrast, no significant differences in total cell binding/uptake or cytotoxicity with 110 nm silver particles were observed between wild-type and SR-A(-/-) macrophages. After administration of 20 nm particles by oropharyngeal aspiration in vivo, wild-type mice and SR-A(-/-) mice demonstrated similar silver clearance from lungs, bronchoalveolar lavage (BAL) fluid, and BAL cells (half-life: 15.5 vs. 23.8 d, respectively) in lungs) after 28 d measured by ICP-MS. However, wild-type mice had increased levels of silver in liver (area under the curve [AUC]: 0.06 vs. 0.02 g·d⁻¹, respectively) and spleen (AUC: 0.16 vs. 0.03 g·d⁻¹, respectively) driven by higher silver levels in liver and spleen at day 7, indicating increased uptake of silver by wild-type mice. After retro-orbital injection, in vivo bioavailability of silver to wild-type and SR-A(-/-) mice were very similar. Overall, these results support the conclusion that SR-A plays a significant role in silver nanoparticle uptake at both the cellular and whole animal levels, but its role may be size-dependent.
and the presence of leukocytes and inflammatory cytokines in bronchoalveolar lavage fluid (BALF). One day after exposure, none of the strains exhibited differences in total protein or LDH activity compared to saline vehicle controls. However, there were marked strain–dependent differences in lung inflammation, with C57BL/6 mice showing a 3-6 fold lower response compared to other strains. Evidence to date indicates that neutrophilic inflammation was attenuated by 24 days, suggesting a transient effect for these nanoparticles. Because C57BL/6 mice were not very responsive, relying on this strain alone to assess NanoAG-induced inflammation may not adequately assess their safety. This work is done in collaboration with the NCNHR Consortium and supported by NIH/NIEHS grants U19ES019545, P30ES07073 and T32ES07032.

1365 Characterization and Toxicity of Silver Nanowires.
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Metal and semiconductor nanowires (NW) have a diverse range of anticipated applications in electrical devices, sensors, catalysts and composite materials but there is very limited information on the fate of NW in the environment or the effect of NW on organisms. We are studying the environmental fate and biological toxicity of silver nanowires (Ag NW), a representative metal NW, to (1) Daphnia magna, an aquatic ecotox indicator; (2) cell lines representing potential target organs, including a cell line from fathead minnow, another eco-indicator species. LC50 values were determined in D. magna exposed to Ag NW and different end-points of cytotoxicity were investigated using cell based imaging assays (high content screening) and other cell assays. Transcriptomic response to Ag NW was also studied in D. magna and in a macrophage cell line using high throughput RNA sequencing. We characterized the physicochemical properties of Ag NW to determine how their chemistry relates to cytotoxicity. We studied the colloidal stability and oxidation rates of Ag NW with two lengths and two coatings in different aqueous solutions, including conditions relevant to studies of toxicity to D. magna, and cell lines. While polyvinylpyrrolidone (PVP) and aluminum (Al) coatings prevent Ag NW aggregation in ultrapure water, rates of aggregation and settling were greatly increased in the presence of dissolved inorganic and organic ions. In stirred anaerobic solutions, morphological changes, including fusing of multiple nanowires, was the dominant transformation. In aerobic solutions, Ag oxidation to release dissolved Ag+ occurred for all coating types. Rates of Ag+ release cannot explain observed differences in Ag NW toxicity to Daphnia, suggesting that toxicity is controlled by a complex mixture of colloidal and chemical processes in solution. Also, in parallel to this physicochemical study, cell internalization, sub-cellular localization as well as in situ localization of Ag NW in Daphnia were carried out.

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Background: Gold nanoparticles (Au-NPs) have been used in previous studies as a particle-specific probe in nanotoxicology studies due to their resistance to oxidative dissolution. Objectives: To examine the particle-specific toxicity of Au-NPs to C. elegans using a toxicogenomic approach and to explore possible mechanisms of defense against toxicity of Au-NPs using gfp strains, mutants and RNAi. Methods: The LC10 of 5.9 mg/L Au-NPs was used for 12hr exposure in microarray experiments using Affymetrix whole genome microarrays. qRT-PCR was used to confirm results for six selected genes. Selected mutant and gfp C. elegans strains were used for toxicity testing to Au-NPs. Differences in distribution of Au in endocytosis mutants was also studied in Daphnia and in a macrophage cell line using high throughput microscopy. Results: 797 significantly differentially expressed genes identified from whole genome microarray analyses were linked to seven common biological pathways. Significant up-regulation was observed for 26 pnp/au genes from non-canonical unfolded protein response (UPR) pathway and molecular chaperones (hsp-16.1, hot-10, hsp-3 and hsp-4). There was significant increase in sensitivity to Au-NPs in non-canonical UPRs (pnp-5 and abu-11) knockouts. Au-NPs induce gfp expression of the vesicular GABA transporter homologue in GABA neurons. Significant up-regulation by Au-NPs of FOXO transcription factor dal-16 was confirmed by microarrays and through using dal-16:gfp strains. Differences in distribution of Au among endocytosis mutants (ehc-1 and rme-2) and wild type nematodes were correlated with differences in toxicity. Conclusions: Our results suggest possible involvement of genes from endocytosis and UPR pathways in uptake and detoxification mechanisms, respectively. Au-NPs may also induce stress response through insulin signaling pathway. Support: US NSF EPA CEINT; University of KY microarray core Pilot grant.

1367 Mouse Neural Progenitor Cells As a Functional Model for Developmental Neurotoxicity Testing Intracellular Calcium Signaling.
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Developmental neurotoxicity (DNT) is often investigated in a traditional manner (in vivo using large numbers of experimental animals), while development of in vitro methods for DNT reduces animal use and increases insight into cellular and molecular mechanisms of DNT. Neural progenitor cells (NPCs) are particularly suited for the investigation of neurodevelopmental processes as they are pre-programmed to differentiate into nervous system-specific cell types: neurons and glia.

The intracellular calcium concentration ([Ca2+]i) plays an essential role in neurotransmission, plasticity and neurodevelopment and can therefore be used as a functional readout for DNT [1]. We therefore investigated changes in [Ca2+]i, evoked by a set of neurotransmitters in primary mouse NPCs using the Ca2+-responsive dye Fura-2. Calcium responses were measured at a single-cell resolution at various differentiation durations. Calcium responses could be evoked by depolarization, acetylcholine, and adenosine triphosphate (ATP), indicating the expression and functionality of the respective neurotransmitter receptors and related calcium signaling pathways.

The data demonstrate that this model allows for the investigation of possible effects of (suspected) developmental neurotoxicants on the development of neuronal characteristics, such as intracellular signaling in response to neurotransmitters. As investigation of chemical-induced effects on the development of functional neuronal characteristics is underrepresented in DNT testing, we argue that biochemical and morphological approaches should be complemented with investigations of neuronal (network) functionality, including network formation, inter- and intracellular signaling and neuronal network function.

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1368 Ketamine-Induced Neuronal Damage and Altered N-methyl-D-aspartate (NMDA) Receptor Function in Rat Primary Forebrain Culture.
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Ketamine, a noncompetitive NMDA receptor antagonist, is used in pediatric general anesthesia and causes neuronal cell death during the brain growth spurt. To understand the underlying mechanisms associated with ketamine-induced neuronal toxicity and search for approaches or agents to prevent such adverse effects, a primary cell culture system was utilized. Neurons harvested from the forebrain of newborn rats were maintained under control conditions or exposed to either ketamine (10 μM), or ketamine plus L-carnitine (1, 30 and 100 μM) for 24 hours, followed by a 24-hour withdrawal period. Ketamine exposure resulted in elevated NMDA receptor (NR1) expression, increased generation of reactive oxygen species (ROS) as indicated by higher levels of 8-oxoguanine production, and enhanced neuronal damage. Co-administration of L-carnitine significantly diminished ROS generation and provided near complete protection of neurons from ketamine-induced cell death. NMDA receptors regulate channels that are highly permeable to calcium. Calcium imaging data demonstrated that neurons exposed to ketamine had a significantly elevated influx of calcium and higher intracellular free calcium concentrations ([Ca2+]i) evoked by NMDA (50 μM), compared to control neurons. These findings suggest that prolonged ketamine exposure produces an increase in NMDA receptor expression (compensatory up-regulation) which allows for a higher/toxic influx of calcium into neurons once ketamine is removed from the system, leading to elevated ROS generation and neuronal cell death. L-carnitine appears to be a promising agent in preventing ketamine toxic effects on neurons at an early developmental stage.
can cause cell death, synaptic remodeling, and altered brain cell morphology. Acetyl-L-carnitine, an antioxidant and neurotrophic dietary supplement, has been reported to prevent neuronal damage from a variety of causes. To evaluate the ability of ALC to protect against propofol-induced neuronal toxicity, neural stem cells were isolated from gestational day 14 rat fetuses and on the 8th day in culture were exposed for 24 h to propofol at 10, 50, 100, 300 and 600 μM, with or without ALC (10 μM). Markers of cellular proliferation (EdU), mitochondrial health (MTT), cell death/damage (LDH) and oxidative damage (8-oxo-dG) were monitored to determine: (1) the effects of propofol on neural stem cell proliferation; (2) the nature of propofol-induced neurotoxicity; (3) the degree of protection afforded by ALC and (4) to provide information regarding possible mechanisms underlying protection. After propofol exposure at a clinically-relevant concentration (50 μM), the number of dividing cells was significantly decreased and oxidative DNA damage was increased. There was also a significant dose-dependent reduction in mitochondrial health as evidenced by decreases in MTT metabolism. No significant effect on LDH release was observed at propofol concentrations up to 100 μM. The oxidative damage at 50 μM propofol was blocked by ALC. Thus, clinically-relevant concentrations of propofol induce dose-dependent adverse effects on rat embryonic neural stem cells by slowing or stopping cell division/proliferation and causing cellular damage. Elevated levels of 8-oxo-dG suggest oxidative damage and ALC effectively blocks at least some of the toxicity of propofol, presumably by scavenging oxidative species and/or reducing their production.

Nicotine treatment resulted in a substantial dose-dependent reduction in mitochondrial function as evidenced by significant decreases in the metabolism of MTT. No significant effect of nicotine on LDH release was observed. 1 μM nicotine significantly increased the expression of activated caspase 3, suggesting that nicotine induced neural stem cell apoptosis. These results suggest that nicotine decreased neural stem cell viability, and nicotine-induced cell death is probably apoptotic in nature.

Cigarette smokers often report having problems associated with disrupted sleep. One potential mechanism underlying sleep disruption could be related to alterations in an individual’s circadian rhythm. Nicotine from cigarettes can activate neuronal nicotinic acetylcholine receptors (nAChRs) throughout the nervous system including structures important for maintaining the circadian rhythm. We have utilized oscillations in larval zebrafish locomotor activity as a model to study circadian rhythms. Initially, the circadian rhythm in embryos reared on a 14/10 light/dark (l/d) cycle was evaluated. Of 257 larvae examined, 76.6% exhibited a typical rhythm where locomotor activity was high in the initial hours of the cycle, declined to a minimum, and then became elevated again at the end of the cycle. We then determined the consequences of raising embryos in constant dark or constant light conditions from 0-127 hours post fertilization (hpf) as well as being exposed to nicotine. At 127 hpf, 23.5% of the larvae reared in constant dark exhibited a typical circadian rhythm and 12.5% of embryos reared in constant light exhibited a typical circadian rhythm. Moreover, the locomotor activity of larvae reared in constant light was reduced and remained at a constant level for the duration of the cycle. Since nAChRs are expressed in the zebrafish pineal gland and retina (two structures that influence circadian rhythms in fish) early in development, we hypothesized that nicotine exposed fish would have a pattern of activity similar to those reared in constant light. Zebrafish reared in 15 μM nicotine from 36-96 hpf and then evaluated at 127 hpf indeed lacked the normal circadian rhythm. The locomotor activity pattern of the nicotine-exposed zebrafish was similar to the pattern of zebrafish reared in constant light; the activity was reduced and remained at constant level for the duration of the cycle. These results indicate that nicotine is capable of disrupting vertebrate circadian rhythms and could be involved in the disrupted sleep pattern of smokers.

In children exposed to alcohol in utero, problems with learning and memory suggest hippocampal involvement and effects on synapse formation and function. Astrocytes may contribute to synaptogenesis by releasing specific factors. The present study investigated how ethanol influences the ability of astrocytes to modulate synapse formation in vitro. Using immunocytochemical labeling of synaptic proteins, confocal imaging and 3-dimensional object analysis, we found that astrocytes pre-treated with ethanol (50 mM) and co-cultured with primary hippocampal neurons (14 days in culture) for 24 hours, induce a 4.5-fold increase in synaptic structure formation. To corroborate that this is reflected in increased functionality, we used whole cell patch clamp techniques to measure spontaneous miniature excitatory post-synaptic currents. In neurons co-cultured with ethanol pre-treated astrocytes, we observed a higher frequency of events, relative to neurons co-cultured with control astrocytes, suggesting more functional synapses in the ethanol group. However, a second population of neurons in the same treatment group was observed to have a lower frequency than control astrocytes. No amplitude differences were observed, suggesting no difference in the number of post-synaptic receptors between groups. Ethanol increases the influx of cholesterol-containing lipoproteins (CCL) from astrocytes, and cholesterol appears to play a role in synapse formation. Direct treatment of hippocampal neurons with high-density lipoproteins induced a 3.2-fold increase in synaptic structures. When cholesterol release from astrocytes was induced by LXR/RXR agonists, through the over-expression of cholesterol transfer protein, this also resulted in an increase in synapses, suggesting a role for astrocytic CCLs in the observed synaptic increase after ethanol pre-treatment (Supported by F31AA019860, AA008154).

PS 1370 Determination of Neurotoxicity in the Developing Rat Induced by Dexmedetomidine.

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Anesthetic, sedative and analgesic drugs are used for diagnostic studies and surgery procedures in infants and children, but little is known about their impact on the developing nervous system. Dexmedetomidine is a selective a2-adrenergic agonist, and has been used in many clinical applications including the reduction of postoperative delirium and has many protective effects on cell damage in animal models. However, dexmedetomidine induced human neutrophil apoptosis by activation of caspases-3, 7, 8 and -9. Based on these, we hypothesize how dexmedetomidine affects neuroapoptosis in the developing rat brain.

Material and Methods: Sprague-Dawley postnatal day 7 (P7) rat pups were used for this study. Rats were divided into three doses of 10, 25, and 50 μg/kg body weight of dexmedetomidine and one control group. Each rat pup received 5 intraperitoneal doses of either saline, or Dex at 90 minutes intervals over 6 hours. After the constant level for the duration of the cycle. These results suggest that nicotine decreased neural stem cell viability, and nicotine-induced cell death is probably apoptotic in nature.

PS 1371 Nicotine-Induced Toxicity in Rat Embryonic Neural Stem Cells.

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Maternal smoking substantially increases the risk of learning disabilities, behavioral problems, and attention deficit/hyperactivity disorder in offspring. Nicotine is the main pharmacologically active component of tobacco smoke. Prenatal exposure to nicotine is capable of causing fetal brain damage. To evaluate nicotine’s effects on the developing nervous system and explore potential mechanisms underlying such toxicity, rat embryonic neural stem cells were used.

Brain cortices were collected from fetal rats (gestational day 14, GD14) for neural stem cell isolation and subsequent culture in commercial rat growth medium. On the 8th day in vitro (DIV), confluent neural stem cells were exposed to nicotine at concentrations of 0.5, 1.0, 2.0, 5.0 and 10 μM for 24 hours. Neural stem cells were identified using monoclonal anti-nestin antibody. Markers of cellular proliferation (EdU), mitochondrial health (MTT), cell death/damage (LDH) and immunohistochemical staining of activated caspase 3 were monitored to determine the nature of nicotine-induced neurotoxicity.
There is credible evidence that environmental factors interact with genetic susceptibilities to determine risk for neurodevelopmental disorders such as autism spectrum disorders (ASD). PCBs and IFNγ are implicated as environmental risk factors for ASD. Increased levels of both have been documented in ASD, and both interfere with neuronal connectivity in central neurons via effects on dendritic growth. Recent reports indicate that PCBs upregulate inflammatory cytokines, but the combined effects of these factors on neuronal connectivity have not been explored. Because of emerging awareness of autonomic dysfunction in ASD, which may link the triad of behavioral deficits, gastrointestinal disturbances, and immune dysfunction often observed in ASD, we are quantifying independent and combined effects of PCB95 and IFNγ on dendritic morphology in cultured sympathetic neurons. Sympathetic neurons were dissociated from superior cervical ganglia of perinatal rat pups and grown in serum-free defined medium in the absence of glial cells. Under these conditions, these neurons do not extend dendrites unless treated with BMP7 (10 ng/ml). After 5 d of exposure to BMPs in the absence or presence of PCBs (1 μM), we observed a selective degeneration of dopaminergic neurons while other types such as GABA neurons, were not affected by dieldrin. Based on these findings, we observed a selective degeneration of dopaminergic neurons while other types such as GABA neurons, were not affected by dieldrin. Based on these findings, we hypothesize that non-dioxin-like (NDL) PCBs mediate this effect. Non-dioxin-like PCBs are widespread environmental pollutants linked to developmental neurotoxicity in children. Recent evidence suggests that non-dioxin-like (NDL) PCBs interfere with neuronal connectivity via interactions with ryanodine receptors (RyRs) which are calcium release channels broadly expressed in the nervous system. NDL PCBs stabilize RyRs in the open configuration and this triggers a calcium-dependent signaling pathway that mediates activity-dependent dendritic growth via CREB activation. CREB activation is also linked to increased formation of dendritic spines and synapses via upregulation of miR132, which suppresses the translation of p250GAP, a negative regulator of synaptogenesis. This suggests the possibility that NDL PCBs modulate these signaling events to influence synaptogenesis. To test this, we quantified dendritic spine and synapse formation in primary dissociated hippocampal cultures and organotypic hippocampal slice cultures treated with PCB 95, a NDL congener with potent activity at the RyR. Nanomolar concentrations of PCB 95 significantly increased spine density and the frequency of miniature excitatory post-synaptic currents. These effects were coincident with upregulation of miR132 and were blocked by inhibiting RyR or CREB, suppressing miR132 or expressing a mutant p250GAP, translation of which is not suppressed by miR132. These data demonstrate that PCB 95 sensitization of the RyR modulates synaptogenesis via activation of a CREB-miR132-p250GAP signaling pathway, and provide further evidence of mechanisms by which NDL PCBs may interfere with normal patterns of connectivity in the developing brain. Supported by NIH grants R01 MH086052, R01 ES014901, P42 ES04693, and the Hope for Depression Research Foundation.

Polychlorinated biphenyls (PCBs) are ubiquitous environmental contaminants that have been linked to behavioral deficits in children and experimental animals. We previously reported that developmental exposure to Aroclor 1254 interferes with spatial memory in weanling rats coincident with decreased experience-dependent dendritic plasticity. We hypothesize that non-dioxin-like (NDL) PCBs mediate this phenomenon via interactions with ryanodine receptor (RyR) Ca2+ channels. To test this hypothesis, we are comparing the effects of developmental exposure to NDL PCB 95, a potent RyR sensitizer, on spatial memory in wildtype (WT) versus mice heterozygous (HET) for the human R163C-RyR1 mutation that enhances sensitivity to haloperidol and haloperidol-like actions. Cultures were exposed to 0, 0.1, 1, or 6 mg PCB 95/kg/day in the diet for 2 weeks prior to mating with Ryr1 WT and Ryr1 HET males, and throughout gestation and lactation. Offspring were weaned at P21 and trained in the Morris water maze from P24-P29. Preliminary analyses show that developmental PCB 95 exposure affects latency, percentage of time (PT) and distance (PD) in the training quadrant, and average distance to the target (ADT) during training and these effects are both sex- and genotype-dependent. Within dose groups, Ryr1 WT weanlings have higher PT and PD and lower ADT than Ryr1 HET littersmates during the probe test. There are no treatment, sex or genotype differences in latency, time in walls, or path length during the visual cue test. There are no differences in dam or pup body weight across experimental groups. These experiments suggest that developmental exposure to a RyR-active NDL PCB congeners alters spatial learning and memory and the magnitude of this effect is influenced by sex and by expression of the R163C-RyR1 mutation. This work was supported by NIH grants R01 ES014901, T32 ES007059 and P42 ES04693 and the ARCS Foundation.

Polychlorinated biphenyls (PCBs) are developmental neurotoxicants and endocrine disrupters that remain widespread in the human food supply. Children exposed during gestation and lactation are at highest risk of neurotoxicity. Genetic differences can influence how these chemicals are metabolized, so we use a mouse model in an effort to identify genes which affect human susceptibility to developmental PCB exposure. Coplanar PCBs activate the aryl hydrocarbon receptor (AhR) in high-affinity Ahrb mice and increase cytochrome P450 (CYP) expression, including CYP1A2 protein which can sequester PCBs in the liver. Our work and the work of others have shown that this sequestration is protective. However, our previous research also found an increase in CYP1A1 mRNA in the brains of our most susceptible mouse line with the AhrCyp1a2(−/−) genotype at postnatal day 28 when PCB tissue levels were highest. Other researchers have reported differential expression and regulation of CYP1A1 and CYP1A2 in the brain. So, we are now assessing CYP1A protein levels and activity in various brain regions. We used the EROD assay as a test for CYP1A1 activity and the MROD assay for CYP1A2 activity. Liver from PCB-treated AhrbCyp1a2(+/+) was used as a positive control whereas Cyp1a1(−/−) and Cyp1a2(−/−) knockout mice served as the respective negative controls. We compared traditional methods used for liver tissue (microsome purification) and a modified EROD method for brain. We also used Cyp1a1 and Cyp1a2(−/−) knockouts in liver microsomes from Cyp1a2(−/−) mice, indicating this is a non-specific test for CYP1A2 activity.

Polychlorinated biphenyls (PCBs) are ubiquitous environmental pollutants linked to developmental neurotoxicity in children. Recent evidence suggests that non-dioxin-like (NDL) PCBs interfere with neuronal connectivity via interactions with ryanodine receptors (RyRs) which are calcium release channels broadly expressed in the nervous system. NDL PCBs stabilize RyRs in the open configuration and this triggers a calcium-dependent signaling pathway that mediates activity-dependent dendritic growth via CREB activation. CREB activation is also linked to increased formation of dendritic spines and synapses via upregulation of miR132, which suppresses the translation of p250GAP, a negative regulator of synaptogenesis. This suggests the possibility that NDL PCBs modulate these signaling events to influence synaptogenesis. To test this, we quantified dendritic spine and synapse formation in primary dissociated hippocampal cultures and organotypic hippocampal slice cultures treated with PCB 95, a NDL congener with potent activity at the RyR. Nanomolar concentrations of PCB 95 significantly increased spine density and the frequency of miniature excitatory post-synaptic currents. These effects were coincident with upregulation of miR132 and were blocked by inhibiting RyR or CREB, suppressing miR132 or expressing a mutant p250GAP, translation of which is not suppressed by miR132. These data demonstrate that PCB 95 sensitization of the RyR modulates synaptogenesis via activation of a CREB-miR132-p250GAP signaling pathway, and provide further evidence of mechanisms by which NDL PCBs may interfere with normal patterns of connectivity in the developing brain. Supported by NIH grants R01 MH086052, R01 ES014901, P42 ES04693, and the Hope for Depression Research Foundation.

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Effect when added to fully differentiated LUHMES or other, non-neuronal cell types. By using an alternative differentiation protocol, LUHMES can be differentiated into neurons without tyrosine hydroxylase (TH), the central enzyme in dopamine synthesis. When exposed during early differentiation (day 0-2), these TH-negative cells were less sensitive to dieldrin compared with normally differentiated LUHMES, indicating that components of the dopaminergic phenotype are responsible for the selective degeneration observed.

1379 Differential Gene Expression in the Neonatal Rat Brain following Combined Maternal Exposure to Chlorpyrifos and Nicotine.

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Environmental exposure to toxic chemicals during early fetal development has been known to manifest long lasting adverse effects for several decades. We hypothesized that combined exposures to pesticides used in households and to nicotine in cigarettes may have adverse synergistic outcome in both the mother and developing fetus. We studied the effects of combined exposure to chlorpyrifos (0.2 mg/kg) and nicotine (1mg / kg) with appropriate controls on the developing rats. Following parturition from exposed mothers, the brains of the new born litters were dissected out to isolate different tissues from the brain. Total RNA from the cerebellum was isolated and global gene expression analysis was done using cell cycle specific super arrays from Qiagen, employing standard molecular techniques. Some of the major findings include the following: A) There were a number of cell cycle genes altered in various treatment groups. B) cdc25b was significantly up-regulated in nicotine treatment group, but was down-regulated in both chlorpyrifos and combined treatment groups. C) cdk5 was up-regulated in chlorpyrifos alone and combined treatment groups. D) Up-regulation of p27k1 and p15ink4b were noted in combined treatment group and not in nicotine or chlorpyrifos alone groups. This data indicates complex pathways of inhibition and acceleration of cell cycle. Further studies with those genes identified as biomarkers from this study can be useful for developing molecular tools for diagnostics and therapeutics for toxicities involving chlorpyrifos and nicotine.

1380 The Effect of Embryonic Exposure to Deltamethrin on the Dopaminergic System and Swim Activity in the Zebrafish.


Pyrethroids are commonly used insecticides that are generally considered to pose little risk to human and environmental health. However, there is increasing concern that children are more susceptible to the adverse effects of pesticides. We developed the zebrafish model to test the hypothesis that developmental exposure to low doses of pyrethroid pesticide deltamethrin alters dopaminergic gene expression and leads to persistent behavioral alterations. Zebrafish embryos were treated with deltamethrin at doses below the LOAEL (0.25-0.5 μg/l), during the embryonic period (3-72hpf) using a static non-renewal water exposure. After 72hpf, embryos were either harvested for RT-qPCR or reared in pesticide free water until adulthood. At 72hpf, deltamethrin treated embryos had significant increased expression of the dopamine transporter (DAT) and dopamine receptors D1 and D2. We quantified the swim activity of 2-week old zebrafish (larvae) and 1-year old zebrafish (adults) using the Noldus Ethovision system and observed significant increases in swim activity in both larvae and male adults that had been developmentally exposed to deltamethrin. To determine if the increased swim activity at the 2-week stage is mediated by deltamethrin’s effect on dopaminergic system, we treated deltamethrin exposed and control larvae to methylamphetamine, a DAT agonist. Methylamphetamine had a significant stimulating effect on control larvae and a significant suppressive effect on larvae developmentally exposed to deltamethrin. To further investigate the role of the DAT, embryos were injected with a DAT morpholino to transiently reduce mature DAT transcript levels during the developmental period. DAT knockdown alone results in significant increases in swim activity at 2-weeks of age which is significantly reduced in deltamethrin treated embryos. Our data suggest that exposure to deltamethrin during the embryonic period results in persistent behavioral changes and is likely due to an interaction between deltamethrin and the dopaminergic system. NIEHS R56ES018863, T32ES007148, R01ES015991

1381 Dysregulation of Redox Homeostasis by Paraquat in Rat Developing Brain.

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Immature brain is susceptible to paraquat (PQ) induced oxidative stress resulting in neuron damage. We hypothesize that dysregulation of glutathione (GSH) homeostasis is a key factor in these neurotoxic responses. To test this, postnatal day 4 (PN4) rat pups were chosen as the period between birth and PN10 includes active brain development equivalent to that in the human 3rd trimester. Initially, LD90 levels on PN4 were evaluated. Sprague Dawley rat pups were grouped as control, vehicle control, and PQ was administered (I.P) at 10, 25 and 50 mg/kg. At 24h post-treatment, declines in viability by 45% and 100% were observed in the 25 and 50 mg/kg PQ groups, respectively. During the 24h period, a decline in motor activity was also noticed in these high dose groups. For subsequent studies the 10 mg/kg dose was used for all time points (2-24 hours). Body weight was unchanged compared to both controls (p<0.05) Brain PQ levels peaked at 4h (1400/mg protein) as assessed by HPLC. In brain, a significant (p<0.05) increase in reactive oxygen levels (ROS) was observed by DCF-DA fluorimetry as early as 6h which persisted to 24h. Increased oxidized form of GSH (GSSG) was observed at all time points (2, 4, 6, 8 and 24h) while decreased reduced glutathione (GSH) was noted only at 24h. These changes were reflected in decreased GSH/GSSG ratios with an associated increase in brain PARP-1 levels (50%) at 24h. These results indicate altered redox homeostasis and DNA damage in PQ exposed brain. The studies have established a role for oxidative stress in PQ induced neurotoxicity in developing brain and a model for studying PQ neurodevelopmental toxicity.

1382 Neonatal Bisphenol A (BPA) Exposure Alters Sex-Specific Estrogen Receptor 2 (ERβ) mRNA Expression in the Postnatal Rat Brain.

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Perinatal life is a critical window for sexually dimorphic brain organization, which is profoundly influenced by endogenous steroids, most notably estrogen. Exposure to endocrine disrupting compounds may disrupt this process, resulting in compromised reproductive physiology and altered sociosexual behavior in adult life. To test the hypothesis that Bisphenol A (BPA) exposure specifically confined to the neonatal critical period impacts ESR expression through weaning, the present study assessed the impact of exposure (on the first three days of life) to 10 μg estradiol benzoate (EB), 50 μg/kg BPA (LBPA) or 50 μg/kg BPA (HBPA) on ERβ2 expression in the bed nucleus of the stria terminalis (BNST), the paraventricular nucleus (PVN) and the anterior medial amygdala (MeA) in the postnatal rat brain. In unexposed animals, ESR2 expression decreased in the BNST and increased in the PVN from postnatal day (PND) 0 to 19 in both sexes. Sexually dimorphic ESR2 expression was transient in the neonatal BNST and PVN, EB decreased ESR2 expression in females in the BNST, and in both sexes in the PVN. In the BNST, ESR2 expression was decreased in males by LBPA and in females by HBPA on PND 10. LBPA increased ESR2 expression in females but decreased it in males on PND 10, thus reversing the sexually dimorphic expression pattern observed in the vehicle controls. In the MeA, BPA mimicked the EB effect and decreased ESR2 expression levels on PND 4. Collectively, these data demonstrate that the neonatal period is vulnerable to BPA exposure, and BPA does not simply mirror the impacts of EB, but rather, disrupted ESR2 expression in a dose, temporal, and region specific manner. The functional significance of this altered ESR2 expression may underlie reported disruptions of adult reproductive deficiencies and abrogated sex differences in sociosexual behavior across the lifespan. Further work will also be needed to establish if these effects can be induced via exposures which recapitulate human exposure conditions and doses.

1383 Effects of Different Endocrine Disruptor Mixtures on Gene Expression in Neonatal Rat Brain Regions: Focus on Developing Excitatory Synapses.

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Brain development is regulated by sex hormones. In mammals, estradiol is thought to control male sexual brain differentiation, but recent data also indicate a role for androgen receptor-mediated mechanisms. Female brain development is dependent
1384 The Effect of Prenatal Exposure to Amorphous Nanosilica Particles on Neonatal Memory.

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Nanomaterials (NMs) are increasingly being used in various fields for their unique functions. Therefore, it is important to ensure the safety of NMs to take advantage of their usefulness. Previously we identified the hazard that intravenous injection of several NMs to pregnant mice might induce intrauterine growth retardation (IUGR). IUGR children are known to have high risks of contracting some diseases, such as neurological disorders. Thus, it is important to assess postnatal effects of prenatal exposure to NMs. In this study, we examined neonatal memory after in utero exposure to amorphous nanosilica particles (nSP), as one of the typical NMs. Pregnant mice were intravenously injected nSP. The body weight of pups were measured weekly. Working memory, reference memory, and context memory of pups were measured by eight-arm radial maze test, Barnes maze test, and fear conditioning test. Pups of nSP treated mice (nSP pups) had smaller body weight than control mice from birth to 3 weeks old. On the other hand, no difference was found on the scores of memory tests between nSP pups and control mice. These results suggest that in utero exposure to nSP might result in growth disorder of pups and have little effect on memory of pups.
Neonatal Exposure to Perfluoroxyhane Sulfonate (PFHxS) in Mice Alters Neuroprotein Levels Essential for the Developing Brain.
I. Lee and H. Vibeberg, Environmental Toxicology, Uppsala University, Uppsala, Sweden.
Perfluoroxyhane sulfonate (PFHxS) is a perfluorinated compound (PFC) used as an industrial additive. The chemical properties of PFCs make them suitable as surfactants and oil- and water repellents, and they are frequently used in products for packaging and as protective coatings. However, the same properties account for their extreme physico-chemical stability, making them practically non-biodegradable, which has generated a worldwide environmental spread and concern. Since we recently have seen that other PFCs, like perfluorooctane sulfonate (PFOS) and perfluorooctanoic acid (PFOA), can induce developmental neurotoxic effects, the purpose of the present study was to explore if neonatal exposure to PFHxS can affect specific neuroprotein levels e.g. calcium/calmodulin-dependent kinase II (CaMKII), growth-associated protein-43 (GAP-43), synaptophysin and tau, in the mouse brain. Male and female NMRI-mice were exposed, on postnatal day 10, to a single oral dose of 6.1 or 9.2 mg PFHxS/kg bw and control animals received a 20% fat emulsion vehicle. The animals were euthanized 24h or 4 months after PFHxS exposure and the neuroprotein levels in hippocampus and cerebral cortex were analyzed. 24h after exposure there were significant differences seen in the neuroprotein levels compared with control and the neurotoxic effects differed between hippocampus and cerebral cortex. Increased levels of CaMKII, synaptophysin and tau in the hippocampus and decreased levels of GAP-43 in the cerebral cortex were measured for both sexes. The effects on the protein levels in adult animals were less pronounced than in the neonates. The results from the present study show that a single oral exposure to PFHxS during a critical period of brain development can cause developmental neurotoxic effects. The neurotoxic effects are similar to previous studies done with other PFCs such as PFOS and PFOA. Further investigations on the developmental neurotoxic effects of PFHxS are needed as there still is very limited knowledge about the neurotoxicity of PFCs.

Prenatal Exposure to 1-Bromopropane Changes Basic Excitability of the Hippocampus and Inhibits the Behaviors Induced by Kainic Acid and Pentyleneetrazole in the Rat Offspring during Lactation Period.
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1-Bromopropane (1-BP), a substitute for specific chlorofluorocarbons, is mainly used for degreasing agents and spray adhesives. 1-BP exhibits central neurotoxicity in adult humans. Animal studies have shown reproductive/developmental toxicity as well as neurotoxicity. However, developmental neurotoxicity remains unclear. We aimed to clarify whether prenatal exposure to 1-BP affects development of neural excitability and behaviors in the juvenile offspring. 1-BP was inhalationally exposed to pregnant Wistar rats from day 1 to 20 (6 h/day) with the concentration of 0, 400, or 700 ppm. On the days of PND 13, 14 and 15, field potentials were recorded from the CA1 area of hippocampal slices obtained from the control and 1-BP groups. Stimulation/response curves of field excitatory postsynaptic potential and those of population spike (PS) enhanced, and the ratios of paired-pulse responses of PS decreased in the 1-BP groups at PND14. PTZ or KA injection to the PND14 offspring caused significant response reduction and increased levels of tau in mice irradiated with 0.5 Gy compared to controls. This demonstrates that a single dose of IR, given at a defined critical period during brain development, is sufficient to cause persistently reduced cognitive functions and increased levels of tau in mice.

CYP1A1_CYP1A2(-/-) Double Knockout Mice Exhibit Impaired Motor Function.
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CYP1A1 and CYP1A2, members of the cytochrome P450 superfamily, are key detoxifying enzymes normally expressed in the liver. Though they are also reportedly expressed in the cortex and cerebellum of the brain, their physiological function in the brain remains unknown. Previous work in our lab uncovered motor deficits in Cyp1a2(-/-) knockout mice. To confirm those findings, we obtained Cyp1a1_1a2 (-/-) double knockout mice which had the Cyp1a2 gene deleted using a different genetic strategy. We compared Cyp1a1_1a2 (-/-) double knockout mice with wild-type Cyp1a1_Cyp1a2 (+/+) mice using a battery of four behavior tests: The rotarod test primarily identifies deficits in cerebellar function related to balance and motor coordination. Gait analysis, adhesive removal, and pole climbing tests primarily identify impairments in the nigrostriatinal pathway, which is a major dopaminergic pathway related to motor function. We found significant impairments in Cyp1a1_1a2 (-/-) double knockout mice in rotarod testing and adhesive removal. Male knockouts demonstrated the greatest impairment in adhesive removal compared with females. Interestingly, Cyp1a1_1a2 (-/-) double knockout mice had shorter latencies in the pole-climbing test. This might be explained by a difference in motivation or anxiety, because the animals climb down a 50cm pole to return to their home cage. The 1-Ahr knockout mice were tested in a gait test for stride length, stride width and stride differential. Together, these data suggest a novel function for CYP1A1 in brain regions important to normal motor function.

Neonatal Exposure to a Single Low Dose of Ionising Radiation Causes Persistent Disruptions in Cognitive Abilities and Increased Levels of Tau in Mice.
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Ionising radiation (IR) is extensively used in the medical field for treatment and for diagnostics. Concern has been raised about possible negative consequences from low dose exposure to IR during critical phases of perinatal and/or neonatal brain development. The brain growth spurt, which is characterized by maturation of axonal and dendritic development, establishment of neural connectivity and acquisition of new motor and sensory abilities, occurs perinatally in humans and neonatally in mice. By using the neonatal mouse as an animal model we are able to study the effect of IR during early periods of brain development and which consequences it has for the adult animal.
Neonatal NMRI mice were irradiated (0; 0.35 or 0.5 Gy) at one single occasion on postnatal day 10. At 2- and 4-months of age, spontaneous behavior was tested in a novel home environment and parameters observed were locomotion, rearing and total activity. Analyses of important neuroproteins in cerebral cortex were performed 24h following irradiation (0 and 0.5 Gy) and at 6-months of age. Observation of spontaneous behavior revealed a significantly deranged behaviour in 2- and 4-month old mice of both sexes irradiated with 0.35 or 0.5 Gy in a dose response related manner. The observed reduced activity during the beginning of the test period and increased activity at the end of the test period indicates a lack of habituation capacity and disrupted cognitive functions. Neuroprotein analyses of cerebral cortex 24h after irradiation and at 6-months of age showed significantly increased level of tau in mice irradiated with 0.5 Gy compared to controls. This demonstrates that a single dose of IR, given at a defined critical period during brain development, is sufficient to cause persistently reduced cognitive functions and increased levels of tau in mice.
cell number, was reduced in PND10, PND21, and PND60 in AhRfx/fx/Math1CRE/+ mice compared to controls, suggesting that cell number was diminished. Following AhR excision, there were fewer proliferating GNPIs in the external germinal layer of the cerebellum at PND10, which resulted in a reduction of granule neurons reaching their final destination in the adult cerebellum. These results suggest that AhR activity plays a role in regulating granule neuron number, possibly by promoting GNP cell cycle activity, which ultimately impacts cerebellar function. These studies will provide novel insights for understanding mechanisms of dioxin-mediated neurotoxicity in the developing cerebellum.

1393 Assessment of Brain Morphometry Data Using Multivariate Statistical Analysis Methodology.
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As required by EPA guidelines, brain morphometry is an area of interest in screening chemicals for developmental neurotoxicity. Brain morphometry data includes multiple measurements from the same animal such as brain weight, brain width, and brain length. These measurements are typically resource intensive and costly and therefore often made on a small subset of animals. Univariate statistical methodology (eg. ANOVA) treats these measurements as independent of one another and analyzes them separately. By contrast, using multivariate analysis of variance, the potential error resulting from conducting multiple univariate tests can be avoided. Multivariate statistical analysis evaluates all measurements simultaneously, taking advantage of the correlation between those measurements. In this simulated study, female brain morphometry data were generated in SAS® using the means, standard deviations, and correlations between measurements observed in an actual study. Three dose groups (control, low, and high) were created. The simulation was designed such that individual brain morphometry components suggested the presence of dose-related increasing trends. A univariate analysis of variance was performed on each of the components: brain weight, brain width, and brain length. The univariate analysis showed no significance for any of the three brain measurements at the 5% significance level. Performing a multivariate analysis of variance yielded an overall significant treatment effect (p < .05). Pairwise comparisons provided evidence that brain morphology for both the low and high dose groups was significantly higher when compared to the control group. In the univariate study, the univariate analysis of variance fails to detect biologically relevant treatment effects in brain morphometry measurements. The multivariate analysis, which addresses the measurements concurrently, identifies statistically significant differences among the dose groups. These results suggest a multivariate analysis should be used to analyze brain morphometry data.

1394 Mother’s Environmental Tobacco Smoke Exposure during Pregnancy and Externalizing Behavior Problems in Children.
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Background: While the impact of active maternal smoking during pregnancy on child health has been well investigated, the association between maternal passive smoking, or environmental tobacco smoke (ETS), and behavioral development of offspring is less clear. This study examines the association between maternal ETS exposure during pregnancy and child behavior problems.

Methods: As part of the Jintan China Cohort Study, mother-child pairs (n=646) were included in the analyses. Maternal ETS exposure at home, the workplace, and other places during pregnancy was retroactively assessed when children were 5-6 years old. Behavior was assessed via the Child Behavior Checklist when children were 5-6 years old. Logistic regression models were constructed to examine associations between maternal exposure to ETS during pregnancy and internalizing and externalizing behavior problems, adjusting for potential confounders including child sex and parental characteristics.

Results: 37% of mothers reported ETS during pregnancy. Children of mothers exposed to ETS during pregnancy had higher scores for externalizing and total behavior problems with 25% of children whose mothers were exposed to ETS, compared to 16% of children of unexposed mothers. After adjusting for potential confounders, ETS exposure was associated with a higher risk of externalizing behavior problems in offspring of exposed mothers (OR=2.08, 95% confidence interval [CI] 1.27-3.43). Analysis after multiple imputations and sensitivity analysis further verified the association, but no dose-response relationship was found. ETS exposure, however, was not associated with internalizing or total behavior problems.

Conclusion: This study suggests that maternal ETS exposure during pregnancy may impact child behavioral development, particularly externalizing behaviors.

1395 Patterns of Clinical Bioindicators in Rat Serum following Acute Exposure to Pesticides of Different Chemical Classes.
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There is interest in bioindicators of adverse outcomes in safety assessment and translational research. Chemically-induced neurological effects may be reflected in specific neuronal changes and/or by general stress-like responses, and such bioindicators may be useful for measuring health impacts. We examined differential profiles of clinical bioindicators after acute treatment (po) with different pesticides (permethrin, deltamethrin, fipronil, imidacloprid, carbaryl, triadimefon). These same rats were evaluated for EEG changes (Lyke et al., Toxicoligist, 2010, 2011, 2012, 2013). Rats were sacrificed after EEG testing. Serum samples were processed by Myriad RBM using their RodentMAP® and Rat MetabolicMAP® assays. Depending on the pesticide, 2-18 of 78 analytes were altered, and each pesticide produced a different pattern of changes. Discriminant analysis indicated that these alterations were best described in a dose response manner for each pesticide. The data show that acute exposure to different classes of pesticides produced different patterns of changes in clinical bioindicators, which may be analyzed for linkages to biological pathways. This is an abstract of a presentation and does not necessarily reflect EPA policy.

1396 Acute Triadimefon-Induced Changes in the EEG of Long-Evans Rats.

We have reported that the non-stimulus driven EEG is altered differently by acute treatment with deltamethrin, permethrin, fipronil, or imidacloprid (Lyke and Herr, Lyke et al., Toxicoligist, 2010, 2011, 2012) in non-restrained animals. In the current study, we examined the ability to detect changes in EEG activity produced by triadimefon, a triazole pesticide which inhibits sterol synthesis in fungal cell membranes and inhibits dopamine re-uptake in the mammalian nervous system. Adult male Long-Evans rats were implanted with epidural screw electrodes. After about 1 week recovery, non-restrained animals were gavaged with corn oil (vehicle) and tested for 2 days for acclimation. On day 3, the rats were dosed with 1 ml/kg corn oil, 75, or 150 mg/kg triadimefon and tested 1 h later. EEG was recorded as 30 segments of 2 s durations, transformed using a Fast Fourier Transformation, and the spectra averaged. These dosages have increased motor activity in Long-Evans rats in previous studies. Treatment with 150 mg/kg triadimefon increased the amplitude associated with Theta activity compared to controls (31% increase), and compared to 75 mg/kg triadimefon (27% increase), when recorded between the visual cortex and cerebellum. These results were different from the decreased gamma activity resulting from treatment with fipronil, increased gamma activity following treatment with permethrin, slightly decreased alpha activity with deltamethrin, and lack of changes after dosing with imidacloprid. The data show that triadimefon can alter CNS activity as measured by EEG, and the alterations may differ from some other pesticides. This is an abstract of a proposed presentation and does not necessarily reflect EPA policy.

1397 In Vitro Assessment of Parkinsonian Neurodegeneration by Dinitrophenolic Herbicides in PC12 Cells.
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Parkinson’s disease (PD) is the most prevalent human neurodegenerative disorder after Alzheimer’s disease affecting 1-2% of the population over 65 years of age. In several epidemiologic studies, chronic exposure to environmental pollutants, in particular pesticides, has been linked to neurodegeneration and the development of PD, though the underlying mechanisms are largely unknown. Research directed by neurologist Dr Peter Nijssen encountered PD patients clustered in a remote rural area in the Netherlands, both occupationally and non-occupationally involved in agriculture, with relatively high exposure to pesticides. Among the pesticides, (dinitrophenolic) herbicides, were identified as common denominators. Since this class of herbicides is known as mitochondrial uncouplers, we hypothesized that they may impact child behavioral development, particularly externalizing behaviors.

Using a battery of in vitro tests, including measurements of cell viability, oxidative stress, single-cell fluorescence calcium imaging and protein analysis, the effects of dinitrophenolic herbicides on different pathways of neurotoxicity were assessed in PC12 cells as a model for mature dopaminergic neurons. Our results demonstrate...
that dinitrophenolic herbicides induce moderate cytotoxicity via calcium-depend¬
ent activation of caspase-mediated apoptosis as well as activation of key proteins character¬
istic for Parkinsonian neurodegeneration in surviving dopaminergic cells. This leads to the hypothesis that human exposure to dinitrophenolic herbicides may be linked to the pathophysiology of PD.

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1398 Protein Cysteine Oxidation and Dopaminergic Cell Death Induced by Pesticides.

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Parkinson’s disease (PD) is characterized by the loss of dopaminergic neurons in the substantia nigra, which is linked to mitochondrial dysfunction, oxidative stress, protein aggregation and impairment in protein degradation pathways. Epidemiological studies suggest that chronic exposure to pesticides is associated with an increased risk of developing PD. However, the underlying mechanisms have not been precisely elucidated. Oxidative protein modifications can modulate the activation of cell death pathways. Thus, we explored the role of oxidative cysteine modifications in dopaminergic cell death induced by pesticides. Human neuroblastoma cells (SK-N-SH and SH-SY5Y) and the fetal human mesencephalic cell line (LUHMES, Lund human mesencephalic) were exposed to paraquat (PQ), rotenone and dieldrin. Compared to neuroblastoma cells, LUHMES cells exhibited greater sensitivity to all pesticides independent from their differentiation state. Apoptotic cell death induced by pesticides was paralleled to oxidative stress and a decrease in glutathione content. Cell death was prevented by glutathione ethyl ester. PQ induced protein alterations in both sulfuric acid (PSSO) and glutathionylation profiles (PSSG). Mass spectrometry of immunoprecipitated PSOHI/PSGG proteins identified several proteins from the ubiquitin/proteasome degradation system and molecular chaperones as being modified by PQSO and SSSG in response to PQ, including E3 ubiquitin-protein ligases (SIAH1, TRIM13), the degradation system and molecular chaperones as being modified by PQSO and SSSG in response to PQ, including E3 ubiquitin-protein ligases (SIAH1, TRIM13, RNF25, RNF139, HERC2, BRE1B), ubiquitin hydrolases (USP47, USP24), ubiquitin conjugating enzymes (UBE20), and Hsp40 co-chaperones (DNAJ4, DNAJC11). The expression of the proteins DNAJC11 was upregulated at a concentration dependent manner by PQ. Results suggest a role for protein cysteine oxidation (sulfenylated and/or glutathionylated) in the regulation of the ubiquitin/proteasome and chaperone-mediated folding systems in response to pesticide exposure. Supported by CONACyT-Mexico (Grant #104316).

1399 Aldehyde Dehydrogenase Dysfunction Increases Parkinson’s Disease Risk via Gene-Environment Interactions.

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Parkinson’s disease (PD) is a neurodegenerative disorder characterized by the loss of dopaminergic neurons in the substantia nigra. We recently reported that exposure to the fungicide benomyl resulted in dopaminergic neuronal loss, aldehyde dehydrogenase (ALDH) inhibition, and an epidemiologic association with increased PD occurrence. Since the dopamine metabolite 3,4-dihydroxyphenylacetaldehyde (i.e., DOPAL) is highly toxic and an ALDH substrate, ALDH inhibition might contribute to the particular vulnerability of dopaminergic neurons. To investigate the potential relevance of this mechanism, we developed a novel ex vivo neuronal assay to screen pesticides for ALDH inhibitory activity. All dichlorodiamine tested (e.g., mancozeb, maneb, ziram), two dichiacarbamates (captan, folpet), and two imidazoles (benomyl, triflumizole) inhibited ALDH activity, potentially via metabolic byproducts (e.g., carbon disulfide, thiophenol). We developed a geographic information system method employing state-mandated commercial pesticide use reports to estimate patient exposures to specific pesticides from 1974-1999. Exposures to ALDH-inhibiting pesticides (i.e., from the screen) were associated with dose-dependent twofold to fourfold increases in PD risk. Patients were geno¬typed for variation in the mitochondrial ALDH2 gene, and haplotype analyses of six single nucleotide polymorphisms revealed that ALDH2 variation potentiated PD risk for people working where ALDH-inhibiting pesticides were sprayed liber¬ally. This ALDH2 model for PD etiology might help explain the selective vulnerabil¬ity of dopaminergic neurons in PD and provide a potential new mechanism through which environmental toxics contribute to PD pathogenesis.

1400 Pesticide-Linked Parkinson’s Disease: Using Drosophila to Build a Novel In Vivo Model of Paraquat, Maneb and Ziram Exposure.

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The vast majority of Parkinson’s disease (PD) cases are sporadic and environmental exposure to pesticides have long been suspected to contribute to PD development, though the disease etiology remains poorly understood. Recently, epidemiological data has supported this and revealed that individual exposures to the pesticides ziram, maneb, and paraquat increase PD risk. Interestingly, it appears that these pesticides interact synergistically and exposure to all three pesticides increases PD risk up to three-fold. We are using the model organism Drosophila melanogaster to study the connection between pesticide exposures and the neuronal dysfunction and dopaminergic cell loss characteristic of PD. We find that combined, but not individual, chronic exposures to maneb and paraquat leads to significant dopaminer¬gic neuron loss. To understand how neuronal dysfunction may contribute to a PD phenotype, we complement this approach by utilizing the Drosophila neuromuscular junction. This powerful tool has allowed us to explore the way pesticide exposures may more subtly affect neuronal physiology and behavior in an intact synapse.

We report that ziram exposure significantly alters synaptic vesicle fusion and rep¬take, processes that are essential for proper neuronal signaling. Ziram has been shown to inhibit the E1 ligase, which is required for targeting proteins to the prote¬asome for degradation. We find that inhibiting the proteasome increases synaptic vesicle fusion in a similar manner to ziram, implying a shared mechanism of action. However, it remains unclear if it is ziram’s inhibition of E1 itself or ziram’s down¬stream effect on the proteasome that is responsible for this synaptic phenotype. Ongoing investigations are exploring these possible neurotoxic mechanisms of ac¬tion and may help elucidate the interrelationship between ziram, neuronal dysfunc¬tion, and early neurotoxic mechanisms that could initiate a PD phenotype.

A number of publications have reported that the i.p. administration of paraquat (PQ) to rodents (normally male C57/B6J mice; up to 3 x 10 mg/kg) results in a loss of dopaminergic neurons from the substantia nigra pars compacta (SNpc), which is the primary area of neuropathological damage in Parkinson’s disease (PD). We have conducted studies in the male C57/B6J mouse examining the effect of high i.p. doses (up to the maximum tolerated dose of 3 x 25 mg/kg) of PQ looking not only for neuronal cell loss, but also for evidence of striatal neurochemical changes and pathological changes using stains to detect disintegrating neurons and neu¬ronal processes and the expected astrocyte/microglial activation in response to neu¬ronal cell damage. Mice were administered paraquat dichloride on 1, 2 or 3 occa¬sions (separated by one week) at doses of 10, 15 or 25 mg/kg. A relatively low dose of N-1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP; 10 mg/kg adminis¬tered i.p. 4 times in a single day at 2 hr intervals) was dosed to a separate group as a positive control. Studies were conducted to GLP and assessment of toxicological endpoints was by individuals blinded to treatment group. No consistent stereo¬logical evidence for a loss of tyrosine hydroxylase positive (TH+) dopaminergic neu¬rones in the SNpc was observed in the PQ treated mice. PQ did not alter the con¬centration of striatal dopamine or its metabolites. Over a range of time points (4-168 hrs post-dose), there was no evidence of PQ-induced neuronal degeneration in the SNpc, degenerating processes in the striatum or apoptotic cell death; astro¬cytes and microglia were not activated. In the MPTP treated mice the number of TH+ dopaminergic neurons in the SNpc was reduced, striatal dopamine was re¬duced and significant pathological changes, including necrotic neurons and astro¬cyte/microglial activation in the SNpc, were consistently observed. These results bring into question the use of PQ mouse studies as a robust model for PD/parkinson¬sonism.
Epidemiological studies have revealed that neural degenerative diseases such as Parkinson disease are typically influenced by genetic and environmental factors, while little is known about the environmental factors. The broad application of paraquat (PQ) has given rise to wide public concern on its potential to damage the nigrostriatal dopaminergic (DA) system because its chemical structure closely resembles that of the well-known DA neural toxicant MPP+. We used male C57BL/6 mice (aged 8 weeks and weight 20 ± 2g) to examine the effects of PQ oral exposure on neural behavior. The Society's criteria for the care and use of laboratory animals were used for all the experiments in this study. After treatment with normal saline (vehicle) and PQ (5mg/kg, 10 mg/kg or 20mg/kg) daily for 30 consecutive days, no significant differences in the body weight gain as well as the brain coefficient were observed among different dose groups. The transmission electron microscope showed that subcellular structures of DA neurons were impaired when exposed to the high dose of PQ. The Open Field test revealed reduced locomotor activity and exploratory drive with the increase of PQ concentrations. Cognitive assessments were conducted in all the groups of mice via Morris Water Maze, no statistically significant difference among treatment groups was shown in the avoidance latency, although a trend toward less learning can be observed. On probe trial, PQ-treated mice did show a significant preference for the correct quadrant. Together, the functional observational battery showed treatment-related, but not necessarily dose-related changes in the mice's reactivity and activity. Thus, we proposed that repeated exposure to PQ can induce neurochemical and persistent neurobehavioral changes.

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the presence of positive cells in the hippocampus of Nrf2-/-- mice treated with 60 mg/kg maneb. Owing to the antioxidant pathway genes included in antioxidant stress and antioxidant defense array, qPCR demonstrated that the only gene mediated by the Nrf2 transcription pathway significantly modulated by at least 1.5-fold was glutathione peroxidase 4 (GPX4). This enzyme was significantly upregulated in Nrf2+/+ mice treated with 30 mg/kg maneb and significantly downregulated in Nrf2-/-- treated with the same concentration. It was concluded that maneb exposure causes oxidative stress mediated by the Nrf2 pathway and that GPX4 plays an important role in this protection.

**1407 Neurotoxicity of the Dithiocarbamate Fungicide Ziram Is Dependent on Synuclein in Zebrafish: Implications for Parkinson's Disease.**

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Parkinson's disease (PD) is a neurodegenerative disorder characterized by dopaminergic (DA) neuronal death and α-synuclein (α-syn) accumulation. Epidemiological studies reveal that exposure to the dithiocarbamate fungicide ziram is associated with increased risk of PD. We have previously found that ziram causes selective toxicity to DA neurons, ubiquitin proteasome system (UPS) inhibition, and α-syn accumulation in primary neuronal cultures. Here, we use developing zebrafish (ZF; Danio rerio) embryos as in vivo model to investigate mechanisms of ziram toxicity and its dependence on endogenous synuclein.

ZF embryos exposed to environmentally relevant levels of ziram were deformed, had altered swimming behavior, and premature death with a LD50 of approx. 50 nM at 7 days post fertilization (dpf). Transgenic ZF expressing green fluorescent protein (GFP) driven by vesicular monoamine transporter protein (VMAT) were used to identify amine neurons in zebrafish. Ziram exposure (50nM) resulted in a 35% reduction in VMAT-GFP-expressing neurons in anterior DA clusters and 26% reduction in posterior clusters (noradrenergic) but was not toxic to non-DA sensory neurons.

ZF express 3 synucleins similar to mammalian β and γ synuclein. Overexpression of ZF γ-syn in neurons led to ZF-syn aggregates and neuronal death in a similar manner as overexpression of human α-syn. ZF embryos exposed to ziram also showed accumulation and aggregation of endogenous ZF γ-syn. This accumulation appeared upon knockdown of ziram’s neurotoxicity since knockdown of ZF γ-syn expression improved survival. Furthermore, treatment with CLR01, a molecular tweezers that mitigates α-syn toxicity, also attenuated ziram toxicity. The mechanism/s by which ziram increases ZF γ-syn appear to involve inhibition of protein degradation since ziram at concentrations as low as 10 nM inhibited the UPS in ZF embryos. These studies add further insight into the mechanism of posttranslational toxicology relevant to the risk of PD.

**1408 Linking Developmental Atrazine Exposure in Zebrafish to Long-Term Neurotransmission Alterations.**

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Atrazine is an agricultural herbicide widely used to prevent pre- and post-emergence of broadleaf and grassy weeds in major crops. It is a major environmental contaminant, commonly present in potable water supplies. Recent work has suggested that atrazine exposure may affect behavior and neurotransmitter systems in rodents. However, rodent studies have typically utilized exposures many magnitudes above environmentally relevant levels, and produced conflicting data. The role of early-life exposures in the development of long-term neurodegenerative diseases has recently received much attention. In an effort to assess the immediate and late-life effects of atrazine on neurological function, zebrafish embryos were exposed to environmentally relevant doses of atrazine (0, 0.3, 3, 30ppb) from 1-72 hours post fertilization. Following the development exposure, larvae were changed to atrazine-free water and allowed to mature under normal conditions to assess lifespan impacts. Neurotransmitter levels were assessed using HPLC with electrochemical detection. Adult female zebrafish (9 months) exhibited statistically significant decreases in 5-hydroxyindoleacetic acid (5-HIAA), the primary metabolite of serotonin, as well as a reduction in 5-HIAA/serotonin turnover. Interestingly, larvae evaluated at 7 days post fertilization did not show significant alterations in neurotransmitters levels, indicating that deficits arise over time, many months after the initial exposure. In addition, microarrays performed on adult female brain tissue revealed alterations in genes that are enriched in neurological, psychological and developmental diseases. Thus, after short-term atrazine exposure during development, persistent neurotransmitter disruptions are observed, indicating long-term disruption of brain function. Future experiments are focused on elucidating mechanisms of neurologic dysfunction and neurodegeneration elicited by toxicant exposure during critical developmental stages.

**1409 Effects of the Long-Term Coexposure to the Herbicide Atrazine and Inorganic Arsenic (As) in the Nigrostriatal Dopaminergic System of the Albino Rat.**

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The herbicide atrazine (ATR) and the metalloid arsenic (As) are substances widely distributed in the environment. In recent years interest has grown regarding the toxicity of these substances on the nigrostriatal dopaminergic system. To date, no study has evaluated the effects of chronic and simultaneous exposure to ATR and As substances that may share the same neuronal target. In order to study the effects of the coexposure As+ATR, six groups of rats were given atrazine (10 mg ATR/kg), arsenic (0.5 or 50 mg As/L of drinking water) or their combinations (ATR+0.5 mg As/L or ATR+50 mg As/L) daily for one year. Behavioral tests showed hypoactivity in the 50 mg As/L group, and hyperactivity in the ATR group. All treatments decreased motor coordination. Striatal DA content was significantly reduced in ATR, 0.5 mg As/L, ATR+0.5 mg As/L and ATR+50 mg As/L groups, compared to controls. The number of mesencephalic TH+ cells was significantly reduced in ATR and ATR+0.5 mg As/L groups, compared to controls. Furthermore, the assessment of cell integrity in the substantia nigra showed decreases in cytochrome oxidase reactivity in all treatment groups, but no changes in malondialdehyde immunoreactivity. In summary, the nigrostriatal dopaminergic system appears to be a target for the toxic effects of ATR and As. Our results indicate that when combined, ATR + As act independently, and ATR has more severe effects on dopaminergic neurons. We appreciate the technical assistance of Biol. Soledad Mendoza. This work was sponsored by PAPIIT 214608-19 and CONACyT 60662, 103907 and 152842.

**1410 Glyphosate Treatment Suggests Offspring and Reproductive Toxicity in Caenorhabditis elegans.**

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Herbicides are widely used in both agricultural and residential areas; therefore, pesticide users, as well as family members, may be routinely exposed to these potentially harmful chemicals. Epidemiological studies have shown pesticide exposure may lead to decreased fertility viability in exposed versus non-exposed females. A significant gap in the literature exists as to whether these observations are applicable to exposures of glyphosate-containing herbicides. To further examine this question, we utilized the model organism, Caenorhabditis elegans, to test our hypothesis that exposure to the glyphosate-containing herbicide, TouchDown (TD), results in decreased numbers of offspring in wild-type (N2) worms. Furthermore, this exposure could render surviving offspring more susceptible to neurodegeneration. Based on concentrations (LC50 and LC75) from previous data, eggs were removed from gravid adults and chronically (24 hours) exposed to TD. The numbers of hatched offspring were counted to determine whether TD adversely affected embryo viability. One-way ANOVA indicated a statistically significant decrease (p < 0.05) in the number of hatched eggs from treated compared to control worms. Our second study was designed to determine whether eggs exposed to TD would result in general neuronal degeneration. Eggs from NW1229 worms (a green fluorescent protein (GFP):neuron construct) were treated as described. Data suggested potential neuronal degeneration in treated worms compared to control (p < 0.05), as determined by fluorescence photomicrograph analysis. Lastly, we chronically exposed eggs from strain CL2166 (GFP::glutathione-S-transferase) to varying concentrations of TD. Fluorescence between control and treated groups, however, did not vary, suggesting that large increases in oxidative stress are not responsible for the observed changes in the nervous system. Taken together, these results suggest that TD decreases the number of viable offspring in the nematode, and that neurodegeneration may be present in those that survive.

**1411 Effects of the Subchronic Exposure to the Herbicide Glyphosate on Behavior and the Nigrostriatal and Mesolimbic Dopaminergic Areas of the Albino Rat.**

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Glyphosate (Gly) is the active ingredient of several herbicide formulations widely used to eliminate weeds. Although it has been described that the Glyph formulations are slightly toxic for mammals, reports of human exposure and studies using animal models suggest that Glyph may share the same neurotoxic targets as other herbicides. In order to evaluate the effects of Glyph on the nervous system, male Sprague-Dawley rats received six i.p injections of 50, 100 or 150 mg Glyph/kg or saline (n=10) during 2
weeks (3 injections/week). We recorded locomotor activity 15 min before and 3 h immediately after each injection. Also, the locomotor activity was recorded during 24 h, 16 days after the last injection. The subchronic exposure to Glyph caused decreases in locomotor activity immediately after each injection, as well as 48 h after the Glyph injection. This hypovigilance was maintained until the 16th day post-treatment. We did not detect changes neither in monoamine concentrations nor in the levels of TPH in NAcc or STR at 2 and 16 days after the last Glyph injection. However, specific binding to D1 dopamine receptors in NAcc decreased dose-dependently when measured 48 h after the last Glyph injection. These results indicate that subchronic exposure to Glyph has acute as well as long-term effects on locomotor activity, and that these effects may be related to a decrement in specific binding to D1 dopamine receptors in NAcc. We thank Soledad Mendroza for her technical assistance.

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1412 Relationship between Administered Dose, Target Tissue Levels and Thermoregulatory Response Alterations after Acute Oral Exposure to the Potent Tremor-Inducing Pyrethroid Bifenthrin in Rats.

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In toxicological studies, potency estimates for pyrethroid insecticides (PYRs) in rats may depend on the exposure and testing conditions. This experimental factor may challenge present efforts to reconsider the health risks posed to humans by exposure to PYRs. We are using exposure-dose-effect studies to explore the influence of the testing conditions on the qualitative toxicological classification of, and experimental potencies for, these insecticides. Four PYRs (tefluthrin, bifenthrin [BIF], deltamethrin, alpha-cypermethrin) are evaluated in Wistar adult rats. Body temperature is measured using microchip-like transponders implanted in rat backs 5-7 days before testing. Subcutaneous temperature (T-SC) is then monitored by radio- telemetry at 30 min intervals for 5 h after oral dosing of PYRs in corn oil. Basal T-SC is recorded before the test day, and 30 min before dosing (i.e., physiological T-SC). The maximal difference in T-SC compared to the pre-dosing baseline (i.e., ΔT-max) is here used as a measure of peak response. Soon after the thermoregulation assay, target tissues are dissected out. Blood, liver and brain (stratum, cortex, cerebellum) are collected at 6 h post-dosing for posterior determination of PYR residues using a SPE-GC-ECD protocol. Results for BIF are shown in this first presentation. 0, 0.5, 3, 9 and 12 mg BIF/Kg (N=4-8) produced dosage-related increases in temperature and ΔT. Dose-related increases in both BIF levels in target tissues and ΔT-max were observed. R-squared values were >0.6 in all pairwise relationshios. These results are mostly consistent with previous BIF studies carried out using motor activity and rectal temperature as endpoints (Wolansky et al., 2006, 2007), and a toxicokinetic study (Scollon et al., 2011).

1413 Interactions of a Promoter/Potentiator of Neuropathy and Estereases of Membrane and Soluble Fractions of Brain.

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Phenylnethyl sulfonyl fluoride (PMSF) is a protease and esterase inhibitor causing protection or potentiation/promotion of the organophosphorus induced delayed neuropathy if dosed before or after the organophosphate inducer. The molecular target/s of potentiation/promotion has not yet been elucidated. The analysis of the protection or potentiation/promotion of the organophosphorus induced delayed neuropathy by PMSF and mipafox at sites other than the substrate catalytic center. Such interactions should be considered to interpret the potentiation/promotion phenomenon of PMSF and to understand the effects of multiple exposure to chemicals.

1414 In Vitro Inhibition and Aging of Neuropathy Target Esterase (NTE) Caused by Fenamiphos and Acetate.

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Organophosphorus-induced delayed neuropathy (OPIDN) is a neurodegenerative disorder characterized by ataxia progressing to paralysis with concomitant central and peripheral distal axonopathy. Symptoms of OPIDN in people include tingling of the hands and feet. These symptoms appear about 8–14 days after exposure. One of the criteria for acceptance of organophosphates pesticides (OPs) in Brazil is if the compound does not cause OPIDN. Guidelines for evaluating OPIDN require the observation of dosed animals over several days and the sacrifice of at least 48 hrs. Adhering to these protocols in tests with analytical standards of OPs is difficult because large quantities of the compound are needed and many animals must be sacrificed. Thus, developing an in vitro screening protocol to evaluate if the OP is able to induce or not delayed neuropathy is important. This study aimed to evaluate in SH-SY5Y human neuroblastoma cell culture samples the potential of some OPs, which are widely used in Brazil, in induce delayed neurotoxicity. The relations between the inhibition and aging of neuropathy target esterase (NTE) by the fenamiphos and acetate were evaluated as indicators of the compound ability to induce OPIDN. Mipafox was used as inducer of OPIDN and paraxoxon was used as non-inducer. The compounds were diluted in ethanol and the cells were incubated with the OPs for 24 hours in a 96-well microplate. Paraaxon exhibited an inhibitory concentration of 50% of enzyme activity (IC50) value approximately 47 times greater than that of the mipafox. Fenamiphos and acetate exhibited IC50 values 117 and 124 times, respectively, greater than that of the mipafox in the SH-SY5Y human neuroblastoma cells. Considering the esterase inhibition and aging results, fenamiphos and acetate would not be expected to have a great ability to induce OPIDN in humans compared with the ability of mipafox.

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1415 Inhibition and Aging of Neuropathy Target Esterase (NTE) in SH-SY5Y Human Neuroblastoma Cells As Screening for Inducers of Delayed Neuropathy.

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Organophosphorus-induced delayed neuropathy (OPIDN) is characterized by a central-peripheral distal axonopathy and Wallerian-type degeneration that develops 8–14 days after poisoning by a neuropathic organophosphorous (OP). The current Organisation for Economic Co-operation and Development (OECD) guidelines for evaluating organophosphorous-induced delayed neuropathy (OPIDN) require the observation of dosed animals over several days and the sacrifice of at least 48 hrs. Adhering to these protocols in tests with analytical standards of OPs is difficult because large quantities of the compound are needed and many animals must be sacrificed. Thus, developing an in vitro screening protocol to evaluate if the OP is able to induce or not delayed neuropathy is important. This study aimed to evaluate in SH-SY5Y human neuroblastoma cell culture samples the potential of some OPs, which are most utilized in Brazil, in induce delayed neurotoxicity. The relation between the inhibition and aging of neuropathy target esterase (NTE) by the trichlorfon was evaluated as a possible indicator of the compound ability to induce OPIDN. Mipafox was used as inducer of OPIDN and paraaxon was used as non-inducer. The compounds were diluted in ethanol and the cells were incubated with the OPs for 24 hours. Paraaxon exhibited an inhibitory concentration of 50% of enzyme activity (IC50) value approximately 47 times greater than that of the mipafox. Trichlorfon exhibited an IC50 value 48 times greater than that of the mipafox in the SH-SY5Y human neuroblastoma cells. Considering the esterase inhibition and aging results, trichlorfon would not be expected to have a great ability to induce OPIDN in humans compared with the ability of mipafox.

1416 Use of a Human Haploid Cell-Based Loss of Functional Genetic Screening Model to Identify Human Susceptibility Genes for Chlorpyrifos Toxicity.

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Chlorpyrifos (CPF), one of the most widely used organophosphorus pesticides, has been known to cause neurotoxicity through the inhibition of cholinesterase activity. However, other neurobehavioral deficits, unrelated to cholinesterase inhibition, have been linked to chlorpyrifos exposure. The mechanisms behind these effects are not fully known or understood. High-throughput loss-of-function genetic
screening tools in yeast or other non-mammalian model systems have been successfully used to identify susceptibility to chemical exposures and decipher chemical compounds mode of action. We recently obtained a newly developed human haploid cell based loss of functional genetic screening model and adopted this haploid cell model in our laboratory. We treated the cells with 200 μM CPF, a dose causing 30% death of haploid cells after 3 days of treatment. After 21 days of treatment, we were able to identify cells carrying genes with deficient functions that play a role in the resistance to CPF-induced toxicity. We are currently conducting functional tests to validate their association with CPF toxicity. Ultimately, this approach will help identify novel susceptibility genes and gain insight into potential mechanisms of CPF-induced toxicity. (This work was supported by Start-up funds to X.R. provide by SUNY Buffalo).

1417 Pharmacological Modulation of Endocannabinoid Signaling Differentially Affects Acute Toxicity of Paraaxon and Chlorpyrifos Oxon in Rats.

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Organophosphorus anticholinesterases (OPs) elicit acute toxicity by inhibiting acetylcholinesterase, increasing acetylcholine levels at cholinergic synapses throughout-out the nervous system. This leads to excessive/prolonged activation of cholinergic receptors and resulting signs of toxicity including increased parasympathetically-mediated secretions (SLUD signs) and involuntary movements (e.g., tremors). Endocannabinoids (eCBs, e.g., anandamide [AEA] and 2-arachidonoyl glycerol [2-AG]) are released from membrane lipids upon neuronal activation. AEA, 2-AG and potentially other putative eCBs can regulate neurotransmission by activating presynaptic CB1 receptors to inhibit neurotransmitter release. We hypothesized that pharmacological modulation of eCB signaling will influence the expression of anti-cholinesterase toxicity. The comparative effects of WIN 55,212-2 (WIN, a CB1 receptor agonist, 1.5 or 5 mg/kg), AM251 (a CB1 receptor antagonist, 3 mg/kg), URB597 (an inhibitor of AEA degradation, 3 mg/kg) and capsazepine (a TRPV1 endovanilloid antagonist, 10 mg/kg) on the toxicity of paraaxon (PO) and chlorpyrifos oxon (CPF) were evaluated. Involuntary movements were induced by both OPs in a dose-related manner (PO: 0.4, 0.5, 1 mg/kg; sc; CPF: 8, 10, 12 mg/kg, sc), whereas somewhat less consistent dose-related effects were noted with SLUD signs. WIN partially reduced signs of toxicity following PO exposure, but had little effect on toxicity following CPO. The CB1 receptor antagonist AM251 had no effect on CPO toxicity, but substantially increased lethality following PO exposure (6/12 vs 1/9). URB597 had no effect on PO toxicity, or on involuntary movements following CPO, but increased SLUD signs following CPO. Similar to effects of URB597, capsazepine had little effect on PO toxicity, but increased SLUD signs following CPO. These results suggest that pharmacological modulation of eCB and/or endovanilloid signaling may differentially influence selected toxic responses to anticholinesterases in an OP-related manner. (Supported by NIEHS R01 ES016308; MPI: WKA & PJL)

1418 Hippocampal Changes Induced by Noncholinergic Disopropylfluorophosphate (DFP) Exposure in Fischer 344 Rat Brain.

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The mechanism of organophosphate (OP) induced inhibition of acetylcholinesterase and subsequent excitotoxicity is well described. However, exposure to OPs at non-cholinergic doses has been reported to cause deficits in cognitive behavior and spatial memory, though little is known about the mechanisms of action. Here an integrated approach using both metabonomic and transcriptomic techniques was used to reveal some of the non-cholinergic effects of DFP in rat hippocampus. Adult male Fischer 344 rats were administered 1 mg/kg DFP or saline (control) via subcutaneous injection at 10 mL/kg. Time points for hippocampal collection were 3, 5, 1, 2, 12, 24 and 48 hr post dose. AChE activity reduced by a minimum of 55% at 2 hr post dose. Total RNA was isolated from hippocampus for differential gene expression analysis using the Affymetrix 1.0 ST gene array at 1 hr post dose. Lipid and aqueous extracts were prepared from each hippocampus at 2 hr post dose, and profiles of small molecule metabolites, lipids and phospholipids were measured using multinuclear NMR spectroscopy. The amino acids valine, isoleucine and alanine were increased 4-5 fold, while co-aminotransaminase and fumarate were increased 2-3 fold after exposure to DFP. Succinate and γ-aminobutyric acid (GABA) were decreased approximately 57%. No change was detected in any of the lipid metabolites measured. Differential gene expression and pathway analysis revealed that the JAK-STAT and the Wnt signaling pathways were down-regulated. By evaluating the impact of low level OP exposure on the metabolic and transcriptomic profile of the hippocampus, we hope to gain a greater understanding of the noncholinergic mechanisms of action and sensitive target areas of OPs.

1419 Dose-Dependent Behavioral Deficits in Egyptian Agricultural Workers with Chronic Organophosphorus Pesticide (OP) Exposures.

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Chronic exposure to OPs is consistently associated with deficits on neurobehavioral tests in workers using pesticides. While years of work have correlated with degree of effect in a few studies, a dose-effect relationship has not been identified, leading some to doubt the association. We identified a population of pesticide application teams in Egypt primarily exposed to one OP, chlorpyrifos (CPF). Teams include engineers (who do not typically enter the fields during applications), technicians (who walk side-by-side with the applicators in the fields), and applicators (who are typically seasonal workers and have the highest exposures). TCPy levels in urine confirmed the pattern of lower to higher exposures across these job categories. Trailmaking, a test of complex visual scanning, motor speed and agility, consists of two test components, A and B that differ in complexity. The test was administered to 54 engineers, 59 technicians, 31 applicators, and 150 control 3 times during the OP application season and 1.5 months after applications had ended. While time to complete Trailmaking A and B improved across sessions, a consistent dose-response relationship was seen in performance speed: Controls had the best (fastest) performance throughout the application season on Trailmaking A (p<0.04) and B (p<0.001). Applicators had slower performance than engineers (p=0.015) and technicians (p=0.032) on Trailmaking A. On the more complex Trailmaking B test, applicators and technicians had comparable performance that was significantly slower (p=0.003 and p=0.012 respectively) than performance of engineers. Test performance at 1.5 months after applications ended revealed that differences between the groups were persistent. Ongoing studies are evaluating relationships between neurobehavioral performance and genetic polymorphisms of enzymes that metabolize CPF. (NIH R01 ES016308; MPI: WKA & PJL)

1420 Chlorpyrifos Exposure and Self-Reported Neurological Symptoms in Adolescent Pesticide Applicators.

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Although studies with adults have demonstrated associations between organophosphorus (OP) pesticide exposure and neurological symptoms, the change in symptom reporting across an application season and the relationship between symptoms and biomarkers of exposure is poorly understood and few studies have examined adolescents. The prevalence of neurological symptoms across the chlorpyrifos (CPF)-application season was examined in adolescent pesticide applicators (n=57) and environmentally exposed adolescents (n=38) in Egypt. Self-reported symptom data at 32 time points over a 7-month CPF-application season were collected. Associations of symptoms with the urine CPF biomarker, trichloro-2-pyridinol (TCPy) and blood cholinesterase inhibition were also investigated. Increased reporting of neurological symptoms were observed among both applicators and non-applicators after several weeks of repeated exposure. Applicators demonstrated a greater percentage of neurological symptoms relative to baseline than the non-applicants at all subsequent time intervals. Cumulative TCPy, an estimate of overall seasonal exposure, was positively associated with average percentage of symptoms only among the applicators in the adjusted models. The prevalence of neurological symptoms among the chlorpyrifos (CPF)-application season was examined in adolescent pesticide applicators (n=57) and environmentally exposed adolescents (n=38) in Egypt. Self-reported symptom data at 32 time points over a 7-month CPF-application season were collected. Associations of symptoms with the urine CPF biomarker, trichloro-2-pyridinol (TCPy) and blood cholinesterase inhibition were also investigated. Increased reporting of neurological symptoms were observed among both applicators and non-applicators after several weeks of repeated exposure. Applicators demonstrated a greater percentage of neurological symptoms relative to baseline than the non-applicants at all subsequent time intervals. Cumulative TCPy, an estimate of overall seasonal exposure, was positively associated with average percentage of symptoms only among the applicators in the adjusted models. No associations were found between acetylcholinesterase (AChE) and butyrylcholinesterase (BChE) inhibition at 1.5 months after applications ended revealed that differences between the groups were persistent. Ongoing studies are evaluating relationships between neu- robehavioral performance and genetic polymorphisms of enzymes that metabolize CPF. (NIH R01 ES016308; MPI: WKA & PJL)
Tetramethylenedisulfotetramine (TMDT) is a highly lethal neuroactive rodenticide responsible for many accidental and intentional poisonings in mainland China. Ease of synthesis, water solubility, potency, and difficulty to treat make TMDT a potential terrorist weapon. We characterized TMDT-induced seizures and mortality in male C57BL/6 mice. TMDT (ip) produced a syndrome of twitches, clenched, and tonic-clonic seizures decreasing in onset latency and increasing in severity with increasing dose; 0.4 mg/kg was 100% lethal. The NMDA antagonist, ketamine (KET, 35 mg/kg) injected ip after the first TMDT-induced seizure, did not reduce lethality, but increased the number of clonic seizures. Doubling the KET dose decreased tonic-clonic seizures and eliminated lethality through a 60 min observation period. Treating mice with another NMDA antagonist, MK-801, 0.5 or 1 mg/kg ip, showed similar effects as low and high doses of KET, respectively, and prevented lethality, converging continuous (status epilepticus) EGG activity to isolated inter-  

dicharial discharges. Treatment with these agents 15 min prior to TMDT administration did not increase their effectiveness. Post-treatment with diazepam (5 mg/kg; GABAA allosteric enhancer) reduced seizure manifestations and prevented lethality 60 min post-TMDT, but icv events were evident in EGG recordings and, hours post-treatment, mice experienced status epilepticus and died. Thus, TMDT is a highly potent and lethal convulsant for which single-dose benzodiazepine treatment is inadequate in managing EEG seizures or lethality. Repeated benzodiazepine dosing or combined application of benzodiazepines and NMDA receptor antagonists are more likely to be effective in treating TMDT poisoning. Supported by NIH NS056093, NS072986, NS044241, DOI PR00634P1 & NYMC.
immunosuppressive agents with harsh side effects. Cannabinoids, which are a group of compounds derived from the marijuana plant (Cannabis sativa), are known to mediate their signals through the cannabinoid receptors, CB1 and CB2, and have been effective as treatment for cancer associated pain, nausea and appetite loss. Recently, their anti-inflammatory properties have been studied. Moreover, the use of cannabinoid therapy for MS has also been exploited. However, the proposed mechanism of action needs to be explored further. We used experimental autoimmune encephalomyelitis (EAE), a murine model of MS, to explore the anti-inflammatory role of cannabidiol (CBD) and its effects on myeloid-derived suppressor cells (MDSCs). EAE disease paradigms were consistently reduced with CBD treatment, as a significant reduction in clinical scores of paralysis and decrease in cellular infiltration, marked improvement of CNS tissue integrity, and reduced demyelination via histopathological analysis were observed. In addition, CBD led to a reduction in the percentage and absolute number of T cells particularly the CD4+ T cells infiltrating the CNS (spinal cord and brain), which were significantly increased in the untreated EAE mice. However, there was a profound increase in MDSs in the spinals, CNS, and peritoneal wash of CBD treated mice as compared to the untreated EAE controls. Both granulocytic and monocytic MDSs were increased in CBD treated EAE mice. Together, these studies demonstrate that CBD treatment may ameliorate EAE via the induction of MDSCs which suppress the aberrant autoimmune response. (Supported by NIH grants R01 AT006688, R01 ES019313, R01 MH094755, P01 AT003961, P20 RR023684 and VA Merit Award BX001357).

1428 Effect of Analgesic Administration on the Guinea Pig Maximization Test.
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Guinea pig maximization tests have been associated with inflammation at cutaneous induction sites due to the use of 1-chloro-2, 4-dinitrobenzene (DNCB) as a positive control and Complete Freund’s Adjuvant (CFA). CFA enhances the sensitization potential of the test substances and its use is required by ISO 10993-10. To alleviate the potential for pain and distress, we evaluated the use of the analgesic, buprenorphine hydrochloride (HCl). Analgesics can modulate the inflammatory response and may interfere with the detection of contact sensitization. The purpose of this study was to determine if the administration of buprenorphine HCl, as a refinement to reduce the potential for pain and distress, would affect the results of the guinea pig maximization test. DNCB and Rubbercoc® Gloves were used as the test articles. The experimental design was consistent with the procedures described in ISO 10993-10 and the guinea pig maximization test (Magnusson and Kligman), with additional parameters evaluated. The experimental design consisted of 10 groups with each group receiving different concentrations of the test articles with or without analgesic treatment. Twenty-four to thirty hours after topical induction, the groups treated with buprenorphine HCl were given 0.006 mg/kg every 12 hours for a total of three doses. Three animals per group were terminated on study day 10 for hematological, coagulation and histological evaluation of treatment sites. The remaining animals were terminated after completion of the sensitization test. Body weight gain, clinical observations, pain assessment, and sensitization potential were evaluated. Clinical observations, hematology, coagulation or histopathology of treatment sites were similar in groups that received analgesics compared to groups that did not. At least 60% to 100% of animals in each group were sensitized with no difference between corresponding groups with or without analgesic treatment. Based upon the results of this study, the use of the analgesic, buprenorphine HCl, did not interfere with the evaluation of the results of the guinea pig maximization test.

1426 Attenuation of Thriceelothene-Mediated Autoimmune Response in iNOS-null MRL+/+ Mice.
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Exposure to trichothelene (TCE), a ubiquitous environmental contaminant, is associated with an autoimmune response both in humans and animal models. However, mechanisms underlying TCE-mediated autoimmunity remain largely unknown. Previous studies from our laboratory in MRL+/+ mice suggest that reactive oxygen and nitrogen species (RONS) may contribute to TCE-induced autoimmunity. The current study was undertaken to further assess the role of oxidative and nitrosative stress in TCE-induced autoimmunity by using iNOS-null MRL+/+ mice. iNOS-null mice were backcrossed to MRL+/+ mice for 10 generations and then N10 heterozygous mutants were intercrossed to obtain homozygous mutants. Female MRL+/+ and iNOS-null MRL+/+ mice were given TCE (10 mmol/kg, i.p., every 4th day) for 6 weeks; their respective controls received corn oil only. TCE treatment led to significant induction of anti-malondialdehyde (MDA)- and anti-4-hydroxynonenal (HNE)-protein adduct antibodies along with increased iNOS in sera, and increased nitrotyrosine (NT) in sera, livers and kidneys in MRL+/+ mice, suggesting an overall increase in oxidative and nitrosative stress. The TCE-induced oxidative stress was also associated with significant increases in serum anti-nuclear antibodies (ANA) and anti-double stranded DNA antibodies (anti-dsDNA) levels. Interestingly, iNOS and NT levels were negligible in both controls and TCE-treated iNOS-null MRL+/+ mice. However, TCE treatment in iNOS-null mice still led to significant increases in serum anti-MDA/ HNE-antibodies along with increases in serum ANA and anti-dsDNA compared to controls. Remarkably, the increase in serum ANA and anti-dsDNA induced by TCE in the iNOS-null mice were significantly less pronounced compared to that in MRL+/+ mice. Our results provide further evidence for a role of oxidative/nitrosative stress in TCE-induced autoimmune response, and iNOS elimination attenuates this autoimmune response. Supported by NIH ES016302.

1427 Mouse Model of Halogenated Platinum Salt Hypersensitivity.
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Occupational exposure to halogenated platinum salts can trigger the development of asthma. Concern for increased asthma risk exists for the general population due to the use of platinum (Pt) in catalytic converters and its emerging use as a diesel fuel additive. To investigate airway responses to Pt, we developed a mouse model of Pt hypersensitivity. Previously, we confirmed the dermal sensitizing potency of ammonium hexachloroplatinate (AHCP) using an ex vivo [3H]methyl thymidine labeling version of the local lymph node assay in BALB/c mice. Here, we investigated the ability of AHCP to induce airway responses in mice sensitized by the dermal route. Mice were sensitized through application of 100 μL 1% AHCP in DMSO to the shaved back on days 0, 5 and 19, and 25 μl to each ear on days 10, 11 and 12. Untreated mice were sensitized with PBS. On day 24, mice were challenged by oropharyngeal aspiration (OPA) with 0 or 100 μg AHCP in saline. Before and immediately after dosing, airway responses were assessed using whole body plethysmography (WBP). A dose dependent increase in immediate airway responses (IAR) was observed in AHCP sensitized and challenged mice. On day 26, changes in ventilatory parameters were analyzed. Mice were exposed to WBP; dose-dependent increases in Mch responsiveness occurred in sensitized mice. Bronchoalveolar lavage fluid harvested from sensitized mice contained an average of 7.5% eosinophils compared to less than 0.5% in control mice (p < 0.05); significant increases in total serum IgE was observed for all sensitized mice. This model will be useful for assessing the relative sensitizing potency and cross-reactivity between different halogenated Pt salts and for investigating the possible adjuvant effects of diesel exhaust particles. This abstract does not represent EPA policy.

Epicutaneous patch tests (EPT) are commonly used to identify chemical agents of allergic contact dermatitis in dermatology patients. Test validity and assessment of allergic reaction severity are highly dependent on the use of reliable chemical allergen test reagents. The purpose of the present study was to measure the actual concentration of nickel sulfate (NiSO4), methyl methacrylate (MM), formaldehyde (FA) and glutaraldehyde (GA) compared to the labeled concentrations of commercial reagents found in dermatology clinics where patch testing is routinely performed. The commercial reagents, NiSO4, MM and GA are supplied either dissolved or suspended in petrolatum (usually in syringe, multiuse containers) while FA is diluted in water. Participating clinics submitted in-date and out-dated reagents to the laboratory for analyses. Both NiSO4 and FA levels were at or above the labeled concentration. NiSO4 particular was uniformly distributed throughout the petrolatum. In contrast, MM was low and variable in commercial allergen reagents. ‘In-use’ MM reagent syringes were all 85% of the 2% label concentration with no observable relationship to expiration date. One MM syringe purchased directly from the manufacturer was 70% of the labeled concentration. Lower MM levels in syringes were consistently measured at the tip vs. plunger end.
of the syringe suggesting loss due to MM’s volatility. GA patch test reagents concentration varied from 27 to 45% of the labeled (1% in petrolatum) amount, independent of expiration date. No GA concentration pattern between tip and plunger was observed. These data suggest that false negative EPT results may be due to instability of volatile or self-polymerizing chemical allergens in the test reagents.

1430 Adjuvant Effect of Dibutyl Phthalate (DBP) in an Animal Model of Contact Hypersensitivity.
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Recent studies have demonstrated that certain phthalates can have adjuvant effects in contact hypersensitivity models, exacerbating inflammatory responses. According to human exposure estimates, perfumes containing DBP can result in topical applications as high as 0.4 mg DPB/day. The aim of the present study was to investigate the adjuvant effect of DBP in the oxazolone-induced contact hypersensitivity model using human relevant doses. Adult male Balb/c mice were divided into 5 different groups (n=6/group). These animals received oxazolone (75 μg/animal) in a hairless abdomen (induction). After five days, mice received oxazolone (75 μg/ear), positive control (DBP-exposed group), or vehicle (negative control group) in the right ear (sensitization). In addition, in the sensitization day and in the two subsequent days, DBP groups received three different doses (0.04, 0.4 and 4 mg DBP/ear) twice a day, while positive and negative controls received vehicle (acetone). All exposures were topical. For three subsequent days after sensitization ear thickness (edema) was measured with the use of a micrometer. After the last measurement, animals were decapitated and the ears were collected for the determination of N-acetyl-fl-D-glucosaminidase (NAG) and Myeloperoxidase (MPO) activity. The study was in accordance to the ethics committee of the Federal University of Paraná. Ear thickness was increased in positive control when compared to vehicle only (negative control) group. No difference was seen between positive control and the lowest DBP dose group. However, oxazolone-induced ear edema was increased in the groups treated with 0.4 and 4 mg DBP/ear when compared to positive control. Similar results were found in MPO and NAG activity. The groups treated with the two highest DBP doses displayed significantly higher enzymatic activity when compared to positive control group. These results indicate that human relevant doses of DBP can have adjuvant effects in the oxazolone-induced contact hypersensitivity in mice.

1431 Characterization of the Mouse Allergy Model to Understand Mechanisms of Drug Allergy.

Developed as a modification of the Lymph Node Proliferation Assay, the mouse allergy model (MAM) appears to be a promising tool for predicting the potential of drug development candidates to produce hypersensitivity reactions (HR). In this model, drugs associated with HR in the clinic produce a marked increase (compared to controls) in the cellularity of the draining lymph nodes (LN). The objective of this study was to characterize the phenotype of draining LN cells to identify new parameters that can be used to enhance the sensitivity and specificity of the MAM and to better understand the mechanism(s) for the response. Drugs that are associated with HR in the clinic (abacavir and amoxicillin) were selected as positive controls for this study. Negative control drugs (merformin and cimetidine) were selected based on the low number of reported HR for these compounds. Groups of 5 mice per group were injected subcutaneously with drug (100 mg/kg) or vehicle once daily for three consecutive days. After a two day rest, cells from the draining brachial LN were isolated and analyzed by flow cytometry. A significant increase in total LN cell number (compared to vehicle) was observed for mice treated with the positive control drugs. Compared to vehicle and negative control animals, an increase in CD4+ and CD8+ T cells and B cells was observed in the draining LN of abacavir and amoxicillin treated animals. Positive control drugs produced significant decreases (~25% compared to control) in the percentage of naïve T cells and increases (~27% compared to control) in the percentage of L-selectin (CD62L) and CD44 double-negative T cells. The negative control drugs induced slight, but statistically insignificant, changes in the expression of these markers. Drugs associated with HR in the clinic produced changes in draining LN cellularity and phenotype that are not observed for negative control drugs. Changes observed in adhesion molecule expression may suggest an effect of positive control drugs on lymphocyte trafficking.

1432 Characterization and Comparison of Methylene Diphenyl Diisocyanate Haptenated Human Serum Albumin and Hemoglobin.
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Methylene diphenyl diisocyanate (MDI) is widely used as a cross-linking agent in the manufacture of polyurethane products. Exposure to isocyanates (dNCO), such as MDI, is known to cause occupational asthma. MDI haptenation of proteins is central to in vitro immunological sensitization, however, the resultant protein conjugates are complex and difficult to characterize. The objective of the present study was to characterize hemoglobin (Hb) and human serum albumin (HSA) following conjugation to different molar concentrations of MDI. MDI-protein conjugates were acid digested to obtain free methylene dianiline (MDA). MDA was extracted, derivatized with fluorescamine and analyzed by HPLC-fluorescence. MDI-Hb was also digested with trypsin and specific amino acid conjugation sites determined by ultra-performance liquid chromatography-quadrupole-tandem time-of-flight mass spectrometry. The trinitrobenzene sulfonic acid assay (TNBS) and denaturing gel electrophoresis were used to determine the extent of cross-linking. MDI conjugation was observed to be dependent on the MDI: protein ratio and the concentration of protein. Greater binding to HSA than Hb was observed and MDI bound to only eight binding sites on Hb compared to twenty for HSA (at 40:1 molar ratio of MDI: protein). Self-polymerization of MDI onto protein was observed on some amino acids at higher MDI concentrations. The TNBS assay was used to confirm cross linking in MDI-HSA with approximately 60% loss of amine reactivity at 0.01 MDI: HSA. More cross-linking was observed at 0.5 mg/ml HSA than at 5mg/ml at 40:1 MDI: HSA. It is concluded that MDI has a greater reactivity toward HSA than Hb with respect to the number of residues haptenated and amount of MDI bound per mole of protein. This work was supported by an Interagency agreement with the NIEHS (YI-ES-0001).

1433 Dimethylfumarate: Potency Prediction and Clinical Experience.
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Dimethylfumarate (DMF) was the cause of a major outbreak of allergic contact dermatitis as a consequence of its use as an antifungal agent in leather products, particularly when used in furniture. This became known, as “toxic sofa dermatitis”. However, what has not previously been established is why the risks to human health had not been assessed adequately for this chemical. The purpose of these investigations was, therefore, to determine whether the frequency and severity of reactions to DMF arose as a function of its intrinsic skin sensitizing potency and/or the nature and extent of exposure. The intrinsic sensitizing potency of DMF was measured using the standard local lymph node assay (LLNA) with determination of an EC3 value; the threshold in the LLNA and which serves as an indicator of relative skin sensitising potency in humans. The EC3 value for DMF was 0.35% when tested in dimethylfumaramide as a vehicle, indicating that this chemical is a strong, but not an extreme, skin sensitizer in this mouse model. DMF was found therefore to have a relative sensitising potency that is comparable with formaldehyde, also a strong human skin sensitizer. The conclusion is drawn that the frequency and intensity of allergic contact dermatitis reactions to DMF suggest that it was the prolonged, repeated and occlusive exposure over large skin areas to this chemical, combined with strong sensitising potency, that together generated the “perfect storm” conditions that caused the DMF epidemic.

1434 Characterization of the Allergenic Potential of Proteins: An Assessment of the Kiwifruit Allergen Actidin.
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Assessment of the potential allergenicity (IgE-inducing properties) of novel proteins is an important component of the overall safety assessment of foods. Resistance to digestion with pepsin is commonly measured to characterize allergenicity, although the association is not absolute. We have shown previously that the measurement of specific IgE antibody production induced by systemic (intraperitoneal) exposure of BALB/c strain mice to a range of proteins correlates with...
Allergic potential. The purpose of the present investigations was to explore further the utility of these approaches using the kiwifruit allergen, actinidin. During the past 10 years, kiwifruit has become an important allergenic foodstuff, coincident with its increased consumption, particularly as a weaning food. The ability of actinidin to stimulate antibody responses has been compared with the reference allergen ovalbumin, and with the non-allergenic protein bovine hemoglobin. Hemoglobin was rapidly digested by pepsin whereas actinidin was resistant unless subjected to prior chemical reduction (conditions more closely reflecting intracellular digestion). Hemoglobin stimulated detectable IgG antibody production at relatively high doses (10%), but failed to stimulate IgE. In contrast, actinidin was both immunogenic and allergic at relatively low doses (0.25% to 1%). Vigorous IgG and IgE antibody production and relatively high titer IgG antibody responses were recorded, similar to those provoked by ovalbumin. Thus, actinidin displays a marked ability to provoke IgE, consistent with allergic potential. These data provide further encouragement that in tandem with analysis of pepsin stability, the assessment of IgE antibody production following systemic (intraperitoneal) exposure of BALB/c strain mice provides an informative approach for the prospective identification of protein allergens.

### 1435 Risk of Allergic Reaction to Undeclared Milk Proteins in a Dietary Supplement.

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In food allergic individuals, the immune system is sensitized to food proteins and mounts a damaging inflammatory response. As a major food allergen, milk and milk-derived ingredients must be declared on food labels under the U.S. Food Allergen Labeling and Consumer Protection Act. Individuals with cow’s milk allergy (CMA) may be at risk from a single exposure to undeclared milk, and may experience hives, swelling, vomiting, diarrhea, and/or asthma. Life-threatening anaphylaxis may occur.

We assessed the risk of allergic reaction due to consumption of undeclared milk proteins in a dietary supplement in adults with CMA by calculating acceptable daily intake (ADI) thresholds for the milk proteins lactoglobulin and casein. ADIs are the lowest exposure that would elicit only mild symptoms in adults with CMA.

We compared these ADIs with estimated intake (EI) levels for the two proteins, based on analytical evaluation of the supplement and recommended serving size.

We conducted benchmark dose (BMD) modeling on data from published double-blind, placebo-controlled milk challenge studies in individuals with CMA. Dichotomous modeling (presence/absence of allergic response) was conducted using EPA’s BMD Software 2.1.2. BMDs and BMDLs (lower 95% confidence interval for the BMD) at a benchmark response (BMR) of 10% for allergic response to milk challenge were modeled.

BMDs were adjusted for milk lactoglobulin and casein content (10% and 83% of total milk protein, respectively) and divided by an UF of 10 (individual variation) to obtain the ADIs. ADIs were the only antibiotics that increased hydrogen peroxide levels compared to the media control at 4h/8h (DNCB, lactic acid, hydrogen peroxide, superoxide), amoxicillin started to increase ROS levels at 4h; these ROS levels were further increased by amoxicillin and cephalexin at 24h compared to the media control. The effects were dose-dependent over a 2-2000μM range. Our data on these canine keratinocytes confirm results published on sulfonamides and human keratinocytes; they also expand them to Blactams.

### 1436 Allergenic Antibiotics (Blactams and Sulfonamides) Induce Oxidative Stress in Keratinocytes.

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Drugs can induce hypersensitivity reactions (drug allergies). Delayed drug hypersensitivity occurs after >5 days of drug exposure. Surprisingly, delayed reactions usually target the skin although most drugs are not administered percutaneously. “Danger” signals such as necrotic cell debris, oxidative stress or inflammation, are key elements in the events leading to the sensitization of the immune system against an environmental toxicant or a therapeutic drug. We hypothesized that allergenic antibiotics would induce oxidative stress in keratinocytes in vitro. We therefore measured levels of intracellular glutathione (reduced and total GSH) and reactive oxygen species (ROS) in a canine keratinocyte cell line (CPEK cells) exposed to 2 Blactams (amoxicillin or cephalexin) 2 sulfonamides (sulfamethoxazole and sulfadimethoxine). Amoxicillin, cephalexin, sulfadimethoxine and sulfamethoxazole decreased levels of GSH in CPEK cells after 24h compared to the negative control; at 8h, levels were similar to the negative control with the 2 Blactams, but higher with the 2 sulfonamides; there was no difference between conditions at 24h. These effects affected total GSH rather than reduced or oxidized GSH. Sulfadimethoxine

### 1437 Activities of Xenobiotic Metabolizing Enzymes in Cell Lines Used for Skin Sensitization Testing In Vitro.

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Skin sensitization is caused by repeated contact with an allergen. An early step in the sensitization process is the interaction of haptons with proteins. In some cases the hapten is formed from pro-haptons by enzymatic reactions in the skin (1). Several in vitro methods are currently in the validation process as alternative methods to test for skin sensitization (2, 3).

The metabolic activities of selected enzymes were assayed in keratinocyte-like cells Keratinosens® (Givaudan, Switzerland) and LuSens (BAFSE, Germany) as well as in the dendritic-like cell U937 (used for the MUST test) and THP-1 (used for the h-CLAT). Protein and cytochrome c reductase contents as well as activities of oxidizing enzymes (CYP, FMO, ADH, AHH), hydrolysing enzymes (esterase) and conjugating enzymes (NAT and UGT) were measured in subcellular fraction of the cells by monitoring metabolic transformation of model substrates. CYP, FMO, UGT and ADH activities were below the detection limit for all investigated cells. NAT and esterase activities were detectable in all cell lines. A1H2 activities were detected in the keratinocyte-like cells Keratinosens® and LuSens but not in U937 or THP-1 cells. These results demonstrate that potential pro-haptons can be metabolically activated during sensitization testing in vitro. The xenobiotic metabolizing enzymes of the respective cells need, however, to be compared to those of native skin.


### 1438 Interleukin 17 Expression by Chemical Allergen-Activated Lymph Node Cells: Selectivity for Contact Allergens.

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Chemical allergens can cause skin sensitization resulting in allergic contact dermatitis, or sensitization of the respiratory tract associated with occupational asthma. The immunological mechanisms resulting in sensitization, and the contributions made by the innate immune response are not fully understood. The interleukin (IL)-17 cytokine family, originally described as Thelper (Th)17 cytokines, are now known to be expressed by cells of the innate immune system. It has been shown previously that topical exposure of mice to 2,4-dinitrochlorobenzene (DNCB), a potent contact allergen, is able to provoke the rapid (within 6-16 hours) production of IL-17 by components of the innate immune system in regional lymph nodes. Here we have examined whether this rapid elaboration of IL-17 cytokines is selective for contact allergens. BALB/c mice were exposed on the dorsum of both ears to various allergens (DNCB, dinitrofluorobenzene, oxazolone, 2-methyl-4-isothiazolin-3-one and 5-chloro-2-methyl-4-isothiazolin-3-one, or α-hexylcinnamaldehyde, to the irritants sodium lauryl sulfate and benzalkonium chloride or to the appropriate vehicle control. At various times thereafter cells from draining lymph nodes were isolated and cultured, and supernatants analyzed for IL-17A, IL-17F, IL-17A/F and IL-22 expression. Expression of IL-17 cytokines and IL-22 was selective for contact allergens with no significant increase in response to skin irritants. A chemical respiratory allergen (trimellitic anhydride; TMA) also induced IL-17 and IL-22, but with substantially delayed kinetics compared with contact allergens. Enhancing the dendritic cell migration kinetics in response to TMA to match those induced by contact allergens did not impact on the tempo of lymph node IL-17 expression. Collectively these data suggest that the rapid elicitation of IL-17 cytokines in draining lymph nodes is a potentially selective biomarker of exposure to contact allergens, and may provide a platform for the development of a novel approach to the characterization of skin sensitizing activity.
Interleukin (IL)-17 cytokines, expressed by T helper (Th)17 cells, play pivotal roles in adaptive immune responses. They have been implicated in autoimmune and allergic diseases and have roles in bacterial and fungal clearance. Importantly, IL-17 is produced not only by adaptive immune cells but also by cells of the innate immune system including the nonconventional γδ T cell subset. It has been shown recently that IL-17 expressing γδ T cells are required for the development of adaptive Th17 responses that mediate experimental autoimmune encephalomyelitis, a murine model of multiple sclerosis. We have examined whether IL-17 influences sensitization to chemical allergens. BALB/c strain mice were exposed topically to the contact allergens 2,4-dinitrochlorobenzene (DNCB), the respiratory allergen trimethyl anhydride (TMA), or to vehicle alone. At selected time points single cell suspensions of draining lymph node cultures were cultured and analyzed for cytokine secretion or for mRNA following enrichment/depletion using magnetic beads. A single exposure to either allergen resulted in transient up-regulation of IL-17 from γδ T cells. Maximal levels of secretion were observed at 6h and 48h following exposure to DNCB and TMA, respectively. After repeated exposure under conditions where DNCB and TMA stimulated polarized Th1 and Th2 cytokine phenotypes, respectively, DNCB, but not TMA, was shown to induce the expression of IL-17. IL-17 production by DNCB-activated cells was shown by complement depletion to reside only in the CD4+ population. In subsequent experiments, responses were explored in γδ T cell null mice and C57BL/6 wild type (WT) controls. A similar pattern of IL-17 production to that provoked in BALB/c strain mice was seen in the WT mice following prolonged exposure to DNCB. However, in the γδ T cell null mice the adaptive Th17 response was completely abrogated. These data suggest strongly that the lack of IL-17 production by γδ T cells during the acute ( innate) response affects the subsequent adaptive Th17 response to contact allergens.

**1440 IL-18 Secretion in Human 3D Organotypic Skin Models for the Identification of Contact Sensitizers.**

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Over the last decade, several in vitro methods have been proposed to identify potential of contact sensitizers, but no accepted in vitro method yet available. Keratinocytes play a key role in all phases of skin sensitization. Interleukin-18 (IL-18) production in keratinocyte was identified as a potentially useful endpoint for determination of contact sensitizers. Limitations of traditional submerged cell culture include chemical solubility, stability in water and metabolic competence. To overcome these problems, experiments were performed to test the possibility to use commercially available 3D organotypic skin models to exploit the possibility to use the secretion of IL-18 for the identification of contact sensitizers. A protocol was developed using different 3D skin models, including EpiDerm TM, EST1000TM. Preliminary experiments were conducted to determine chemical toxicity profiles using the MTT viability assay. Additional doses were then chosen for IL-18 secretion experiments. Following topical exposure for 24 h to several contact allergens (including benzocaine, cinnamaldehyde, cirtin, DNCB, eugenol, isoeugenol, 2-mercaptobenzothiazole, oxazoline, etc.) and irritants (including chlorobenzene, lactic acid, octanoic acid, phenol, salicylic acid, SDS, Tween 20, etc.) a robust in vitro IL-18 secretion to either allergen resulted in transient up-regulation of IL-17 from γδ T cells. Maximal levels of secretion were observed at 6h and 48h following exposure to DNCB and TMA, respectively. After repeated exposure under conditions where DNCB and TMA stimulated polarized Th1 and Th2 cytokine phenotypes, respectively, DNCB, but not TMA, was shown to induce the expression of IL-17. IL-17 production by DNCB-activated cells was shown by complement depletion to reside only in the CD4+ population. In subsequent experiments, responses were explored in γδ T cell null mice and C57BL/6 wild type (WT) controls. A similar pattern of IL-17 production to that provoked in BALB/c strain mice was seen in the WT mice following prolonged exposure to DNCB. However, in the γδ T cell null mice the adaptive Th17 response was completely abrogated. These data suggest strongly that the lack of IL-17 production by γδ T cells during the acute ( innate) response affects the subsequent adaptive Th17 response to contact allergens.

**1441 Animal Strains Usable to the LLNA: brdU-ELISA (OECD 442B).**

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The murine local lymph node assay (LLNA) is a widely accepted alternative test to assess the skin sensitizing potential of chemical substances. The original LLNA was designed to quantify lymph cell proliferation in the draining auricular lymph nodes by incorporation of radioabeled thymidine analogues. To avoid use of the radioisotope and to avoid some of the limitations it poses, several modified LLNA protocols were developed. One of the modified methods utilizes the nucleoside (uridine) analogue of 5-Bromo-2-deoxyuridine (BrdU). In this method, the incorporated BrdU is measured by ELISA, using anti BrdU antibody. This modified LLNA:brdU-ELISA has been validated and adopted as OECD test guideline 442B. In this guideline, the CBAj or CBA JN strains are recommended, since research strains during the validation. Previously, we noted differences in response in the LLNA with different strains of mouse. In this study, we used mice of several CBA strains to compare the reactivity against several skin sensitizers and a non sensitizer. It is confirmed that the mouse strains, CBAj/ColaHsd and CBA/N Slc showed similar but not equal response to CBAj/NglJ, the preferred strain in the LLNA:brdU-ELISA.

**1442 Polarized Immune Responses Induced by Chemical Allergens Display Differential DNA Methylation Patterns.**

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There is increasing evidence that epigenetic regulation of gene expression plays a pivotal role in the orchestration of immune responses. It has been demonstrated previously that chemical allergens can be divided into two categories: contact allergens (such as dinitrochlorobenzene; DNCB), that cause type 1/type 17 polarized responses and respiratory allergens (such as trimellitic anhydride, TMA) that induce a preferential type 2 response. Such polarization occurs upon repeated (13 day) topical exposure. In order to explore the regulation and maintenance of such responses at the molecular level, the genome wide pattern of DNA methylation following treatment with the reference allergens DNCB and TMA was characterized. Mice (n=5 per group) were exposed to DNCB, TMA or vehicle alone for 13 days, and draining auricular lymph nodes excised. DNA was extracted, sonicated and processed for methylated DNA immunoprecipitation (MeDIP) followed by hybridization to a high resolution DNA promoter array (Roche) representing 28,000 probe ranges, covering promoter regions and known CpG islands. Changes in DNA methylation profiles for allergen-activated tissues were compared with the vehicle-treated control samples, with a cut off of p<0.01. More differentially methylated regions (DMR) were recorded for DNCB tissue than for TMA tissue (6319 versus 2178), although approximately half those identified for TMA were in common with DNCB. Direct comparisons between DNCB and TMA-treated tissue revealed 268 DMR that were differentially regulated between the two different classes.
of allergens. The 15 most significant genes identified were associated with hypermethylation and hypomethylation for DNBCB and TMA exposure, respectively. Thus, topical exposure to different classes of chemical allergen under polarizing conditions results in characteristic patterns of DNA methylation indicative of epigenetic regulation of the developing allergic response. Furthermore, DNBCB and TMA are associated with more silenced and activating epigenic marks, respectively.

Developmental and reproductive toxic effects of some naturally occurring Fusarium mycotoxins were shown in certain animal models. In this study we used the nematode Caenorhabditis elegans (C. elegans), as an alternative model for mechanistic studies, to investigate the developmental and reproductive toxic effects of T-2 toxin (T-2), zearalenone (ZEN), and deoxynivalenone (DON). C. elegans (N2) 1-day or 3-day old were exposed to various concentrations of these toxins for up to 72 hrs. The development arrested rate (DAR) and the reproduction arrested rate (RAR) were outcomes used to assess the overall effects on the development and reproduction of the exposed C. elegans. T-2 has the most potent suppressive effects on the development and growth, concentrations from 0.5- to 8-mg/L caused averaged DAR from 12.01% to 41.6%; exposure to ZEN from 5- to 80- mg/L caused averaged DAR from 11.14% to 35.56%; exposure to DON from 50- to 800-mg/L caused averaged DAR from 12.71% to 32.77%. The concentration ratio for inducing significant DAR (10%) is approximate to 1:10:100 for T-2, ZEN, and DON, respectively. T-2 also has the most potent toxic effect on the reproduction, concentrations from 0.5- to 8-mg/L caused averaged RAR from 28.97% to 72.43% with an EC50 at 3.57-mg/L; exposure to ZEN from 5- to 80- mg/L caused averaged RAR from 26.15% to 90% with an EC50 at 12.02-mg/L; exposure to DON from 50- to 800-mg/L caused averaged RAR from 23.83 to 52.33% with an EC50 at 585.85-mg/L. The concentration ratio for EC50 is approximate to 0.13:1:64.1 for T-2, ZEN, and DON, respectively. These results suggest that T-2 and ZEN have more significant toxic effects than DON on development and reproduction in C. elegans and that C. elegans is an excellent model, due to its short life cycle and easy genetic manipulation of genes, for studying molecular mechanisms of developmental and reproductive toxic effects caused by food-borne mycotoxins.

1447 Zebrafish As a Complementary Model in Toxicology.
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Growing awareness to apply the principles of Replacement, Refinement and Reduction (3Rs) of animals in regulatory testing drives the need for alternatives identifying potential toxic agents with accuracy, speed, reliability and respect for animal welfare. So far, for complex endpoints like reproduction and developmental toxicology, animal-free in vitro models are limited and cover only a restricted part of the reproductive cycle. Various characteristics warrant the Zebrafish (embryos and/or larvae) as an ideal non-mammalian whole organism model that could bridge gaps between in vitro cell systems and complex reproduction studies in mammals, e.g. small size, ease of obtaining high number of progeny, external fertilization, transparency and rapid development of embryo, and a basic understanding of its gene function and physiology.
Macroscopic examination of zebrafish embryos/larvae predicted toxicity of chemical and pharmaceutical agents with high certainty and proved to form a reliable total organism approach to study embryo- and developmental (neuro)toxicity. More in depth analysis and interpretation of the locomotor data in the developing zebrafish larvae relative to (nervous system) development increases the use of the zebrafish model in developmental neurotoxicity research. A full histopathological survey of the embryo’s combined with high-end analytical methods appeared to further increase the applicability domain of zebrafish into toxicity screening studies. Further refinement with respect to the physical and chemical properties of test compounds that deter the body burden and tissue distribution increases the predictability of the assay. For translation to other toxicity models and man, there is an urgent need to research on responsible toxicity pathways. This research is supported by the Dutch Ministry of Health, Welfare and Sport and the Dutch Ministry of Social Affairs and Employment.
strains were exposed in triplicate for five and fifteen generations to the IC20, 50% IC20, and 25% IC20 dieldrin OCP concentrations. The oligonucleotides unique to each deletion strain were PCR-amplified and hybridized to TAG arrays to identify sensitive, unaffected, and resistant strains. The overrepresented biological terms within the data assisted in the selection of individual deletion strains for confirmation experiments. Analyses indicate that amino acid sensing and components of the pyruvate dehydrogenase complex are critical for cell survival under dieldrin exposure. Exogenous amino acids rescue dieldrin toxicity, while lower concentrations of amino acids in the media exacerbate toxicity. Finally, it was demonstrated that dieldrin inhibits amino acid uptake in yeast cells. Future investigations will examine whether amino acid status is tied to dieldrin toxicity in human cell lines. Additionally, the development of an automated high-throughput system to screen the yeast deletion library for altered growth in various toxicants may be discussed.

1449 Evaluation of Epigenetic and Developmental Effects in Danio rerio Zebrafish Embryos following Diethylstilbestrol, 17β-Estradiol And/or 5-AzaCytidine Treatment. R. G. Ellis-Hutchings, R. A. Alvey, A. V. Marshall, M. J. LeBaron, R. Sura, N. P. Moore, B. Gollapudi, and R. J. Rasoulpour. Toxicology and Environmental Research & Consulting, The Dow Chemical Company, Midland, MI. The fundamental mechanisms that regulate epigenetic modifications as well as embryonic development are highly conserved amongst vertebrates. Danio rerio (zebrafish) potentially may be used to examine epigenetic alterations during stages of early development. Zebrafish embryos were treated with estrogen molecules (17β-estradiol (E2) or diethylstilbestrol (DES) (0.001, 0.01, 0.1, 1, 15, 2, 2.5, 3, 5, 7, and 10μM) in the presence or absence of a DNA methyltransferase inhibitor 5-azaCytidine (5-aza-C, 0–100μM) and evaluated for viability. Only strain 5, a developmental abnormality was assessed using a quantitative scoring system: somites, notochord, tail, fins, heart, face, brain, arches and jaw. Treatment with 25μM 5-azaC prior to the period of DNA re-methylation (2-cell stage) caused >80% lethality whereas starting treatment shortly after the onset of DNA re-methylation (a 16-cell stage) resulted in <10% lethality. Regardless of developmental stage, treatment with 50 and 100μM 5-azaC caused >90% lethality. Treatment with >2μM DES caused significant lethality whereas treatment with 1 or 1.5μM increased the incidence of developmental variations. Treatment with >7μM E2 significantly impacted survival, while 5μM caused a generalized slight increase in developmental abnormalities; however, 1, 1.5, 2, 2.5, and 3μM E2 caused slight developmental effects on the heart, face and brain. Overall, E2 had a wide developmental dose response curve while DES caused significant lethality within a narrow region. Morphological characterization of embryos and larvae following exposure to these compounds indicated that within the chosen dose response range, clear apical end point not observed-adverse-effect levels (NO(A)ELs) and lowest-observed-adverse-effect levels (LO(A)ELs) can be determined thereby providing a useful model to establish associations between apical and epigenetic end points.

1450 A High-Throughput Mechanism-Based Toxicity Screen Using C. elegans. R. B. Goldsmith1, J. R. Pirone1, W. A. Boyd1, M. V. Smith2, and J. H. Freedman1,3. 1NIH, Research Triangle Park, NC; 2NTP, NIEHS, Research Triangle Park, NC; 3SRA International, Durham, NC. To quantitatively assess the effects of chemical exposures on transcription, an automated high-throughput in vivo toxicity assay was developed to measure changes in the levels and cell-specificity of specific gene expression in C. elegans. Transcriptional responses were measured in individual strains of transgenic C. elegans that express fluorescent proteins under the control of archetypal stress-inducible genes: cel-3, esp-35K2, gpr-1, gpr-38, gor-4, hsp-16.2, hsp-16.41, hsp-17, hsp-4, hsp-6, hsp-60, mtl-2, ugt-1 and ugt-13. As part of assay development, transgenic nematodes with large dynamic ranges (i.e., low constitutive levels of expression) contained a concentration-re- 

desc and sensitivity (high response at low concentrations) were identified following exposure to juglone, an oxidative stressor; N-methyl-N’-nitro-N-nitrosoguanidine, a DNA damaging agent; cadmium, a heavy metal; chlorpyrifos, an organophosphate neurotoxin; tunicamycin, an endoplasmic reticulum stressor; and heat shock, a protein denaturant. Concomitant with chemical exposure, fluorescence data were measured from images acquired using a high content imager and subsequently analyzed using CellProfiler’s WormToolbox, a high-throughput imaging analysis program designed for use with nematodes. For some of the stress-inducible genes high levels of constitutive expression limited their dynamic range and utility in this assay. As expected, not all of the chemicals affected all of the genes; however, the chemicals generally affected the stress-inducible genes that corresponded to their known mechanisms of action. For example, nematodes containing the hsp-1 transgene, which responds to endoplasmic reticulum stress, showed increased fluorescence after tunicamycin treatment. In addition, strong correlations were observed between chemical concentration and fluorescent signal. The results obtained using the transgenic C. elegans were verified by measuring changes in the cognate steady-state mRNA levels by qRT-PCR. These results show that this assay can be used to assess the toxicity of genes on expression in vivo.

1451 The Effects of Microcystin-LR on the Chemotaxis Indexes of Caenorhabditis elegans. C. Moore and B. Puchner. Molecular Biosciences, University of California Davis School of Veterinary Medicine, Davis, CA. The blue green algae toxins microcystins (MCs) are known ubiquitous hepatotoxins found in lake and marine waters. The World Health Organization has established a drinking water guideline of 1 μg/L total MCs based on MC-LR liver toxicity data after oral exposure. MCs inhibit serine/threonine protein phosphatases (PP1 and 2A), a mechanism extensively studied in the liver; yet the effects of MCs on the nervous system are poorly understood. To study MCs’ effects on the nervous system, a 24-hour exposure assay using the neurotoxicity model Caenorhabditis elegans (C. elegans) was established. C. elegans have remarkable genetic and neuro-biochemical conservation with humans, and with all 302 neurons extensively studied, key sensory neurons have been linked to specific behaviors. It is well known the AWA and AWC sensory neurons detect diacetylated benzaldehyde, respectively, and sensory neuron function can be quantified using a chemotaxis index (CI). Wildtype (N2) adult nematodes were exposed for 24 hours to 0, 1, 10, 40, 80, 160 and 320 μg/L MC-LR before determining CIs. No significant changes in CIs for benzaldehyde were observed. At MC-LR concentrations of 40 μg/L and above, CIs for diacetyl decreased significantly. The unchanged CIs for benzaldehyde suggest that AWA is not affected. The decreased CIs for diacetyl suggest that AWA is targeted by MC-LR. PP2A inhibitor okadaic acid and PP1 inhibitor tuamotycin were used in the same 24-hour exposure model to evaluate whether PP1 or PP2A inhibition result in changes in AWA function. Results of CI indexes suggest that the AWA chemosensory neuron is affected by MC-LR resulting in neurotoxicity.

1452 Quantitative Assessment of Phase II ToxCast™ Chemical Toxicity in C. elegans. W. A. Boyd1, M. V. Smith2, J. R. Pirone2, J. R. Rice1 and J. H. Freedman1,3. 1DNIEHS, NTP, Research Triangle Park, NC; 2SRA International, Research Triangle Park, NC. 3EPA, NIEHS, NTP, Research Triangle Park, NC. The Tox21 community is working to prioritize thousands of chemicals for further toxicity testing and to develop prediction models for human toxicity. To integrate the large amounts of high throughput in vitro and in vivo toxicity and develop these models, better quantitative methods are necessary. To develop quantitative assessments of in vivo toxicity data, the U.S. EPA’s ToxCast Phase II library, which contains 676 unique chemicals including failed drugs, food additives and industrial products, was screened using the C. elegans growth assay. This assay uses the COPAS Biosort to measure size changes in individual nematodes after 48-h exposures. Over the seven concentrations tested (0.5–200 μM), >50% of the chemicals caused a decrease in growth. To rank their toxicity in C. elegans, a concentration-response curve for each chemical was described using isotonic regression and the statistical significance of the curve assessed. For significant concentration-response curves, the fitted isotonic regression model calculated three parameters that characterize the efficacy and potency of each chemical: 1) change in response between control and highest concentration (R); 2) concentration of chemical at which half of R is reached (ECR50); and 3) slope of the curve at the ECR50. Chemicals were also ranked by computing a toxicity score, which is a weighted sum of the change in response at each dose relative to the control. The top 5% most active compounds included several organic pollutants (e.g., DDT, PFOS), which have been banned from use due to their toxicity and bioaccumulation. Excellent reproducibility was observed for eight chemicals replicated in the library; seven were active and one was inactive for all replicates. The application of this new method for the quantitative assessment of chemical toxicity in C. elegans could be applied to other in vitro and in vivo toxicity screens, thus allowing for better comparisons among various assays, and the development of predictive models of toxicity.

1453 Triazole-Induced Gene Expression Changes in the Zebrafish Embryo. S. A. Herrmsen1,2, T. E. Pronk3, E. van den Brandhof1, L. T. van der Ven1 and A. H. Piersma1,3. 1Health Protection, RIVM, Bilthoven, Netherlands; 2Toxicogenomics, University, Maastricht, Netherlands; 3IRAS, University, Utrecht, Netherlands. Sponsor: H. van Loeveren. The zebrafish embryo is a promising alternative test for developmental toxicity. The classical read-out is via morphological assessment. Microarray analyses may increase sensitivity and predictability of the test by detecting more subtle and mechanistic
responses. We have shown earlier that with transcriptomics data we could discriminate between two chemical classes, glycol ethers and triazoles, in a concentration-response design. It is time-consuming and expensive to perform concentration-response analysis for many compounds. Thus, we studied the possibility of relating gene expression profiles of structurally related chemicals tested in a single concentration, to a complete transcriptomic concentration-response of the triazole anti-fungal fluazolamide (FLU). We tested five other triazoles, hexaconazole (HEX), cyproconazole (CYP), triadimefon (TDF), myclobutanil (MYC), and triticonazole (TTA) at equipotent concentrations based on morphological evaluation. Results showed that compounds differed in their regulation of major anti-fungal and developmental toxicity pathways, steroid biosynthesis and retinol metabolism, respectively. Assuming that the ratio between these pathways is relevant for efficacy compared to developmental toxicity, we found TCT was more efficacious and CYP was more toxic compared to the other triazoles. MYC showed a different response similar to the high toxic concentrations of FLU. We here demonstrated that gene expression data allow more comprehensive assessment of compound effects by discriminating relative potencies using these specific gene sets. The zebrafish embryo model can therefore provide information on relative pathway sensitivity related to intended mechanism of action and toxicological activity of compounds.

1454 Early-Life Exposure to Methylmercury and Multiple Stressors: Daphnia pulex As an Alternative Model System to Evaluate Long-Term Effects.

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While human populations are exposed simultaneously to chemical stressors and physical factors including temperature and differing nutrition, safe limits of exposure are based primarily on tests using single chemicals. Due to the high cost and time required to test mixtures in traditional animal models, development of novel model systems is of critical importance. We are currently evaluating Daphnia pulex, a standardized USEPA and OECD ecotoxicology model system and an NIH model organism for biomedical research, as a tool for assessing long-term effects of early life exposures to multiple stressors. As a case study, we are examining the interaction between MeHg exposure and varying temperature regimes and nutrition. We hypothesize that: Early life exposures in high temperature or a low food regime will increase MeHg toxicity as measured by reproduction and lifespan in Daphnia pulex. D. pulex were exposed to a matrix of three temperature regimes emulating standard laboratory conditions versus natural environment daily fluctuations, three feed rates and five concentrations (200ng/l to 1000ng/l) of MeHg as methylmercury (II) chloride. Data evaluating lifespan and fecundity across the temperature regimes suggest that although there are no significant differences in time to first reproduction, D. pulex lifespan and fecundity were increased in daily fluctuating versus standard laboratory conditions and those that would be experienced in a natural environment. Subsequent work will evaluate interactions. Since current alternative model systems are not designed to evaluate long-term effects of early life exposures, our research is designed to evaluate the utility of D. pulex as a complementary model system.

1455 A Rapid Chemical Screening Platform in C. elegans for Assessing Environmental Germline Disruption.

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Despite the developmental impact of errors in chromosome segregation, we lack the tools to comprehensively assess environmental effects on meiotic integrity in animals. Here, we report the development of an assay in C. elegans that fluorescently marks aneuploid embryos following chemical exposure. We qualified the predictivity of the assay against chemotherapeutic agents as reference compounds, as well as environmental compounds with comprehensive mammalian in vivo endpoint data. The assay was highly predictive of mammalian reproductive toxicities with a maximum specificity of 78%. Finally, we validated selected compounds from the screen by analyzing germline maintenance following exposure. With this novel approach, we provide the first high-throughput screening strategy for the assessment of environmental effects on the germline.

1456 Cytotoxic Effects of Methyl-Mercury in Whole Worms and Pan-Neuronal GFP-Expressing Embryonic Cultures of Caenorhabditis elegans.

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Caenorhabditis elegans (C. elegans) strain KCl136 was used to study whole worm viability in response to methylmercury (MeHg) toxicity in a semifluid gellan gum medium. Stage L3 KCl136 whole worm viability is reduced by exposure to 0.5-25 μM MeHg (p=0.0001) for 24-48 hrs. At the highest [MeHg] tested, cytotoxicity was 54% and time of exposure (24 and 48 hrs) did not have an effect (p=0.15). C. elegans strain NW1229, which expresses pan-neuronal Green Fluorescent Protein (GFP), was used to prepare primary cell cultures from eggs to test the ability of the nematidine-A to reduce the cytotoxicity of MeHg in worm neurons. Nematidine-A is a novel dihydropyridine (DHP) L-type voltage gated Ca2+ channel antagonist and we hypothesize it has a similar effect as we have previously reported for the non-DHP-type antagonist verapamil. At [MeHg] of 0.4-1.2 μM, cytotoxicity in neurons was both concentration and time dependent as determined by ethidium homodimer viability assay. Thus the results obtained in whole worms differ somewhat from those isolated neurons relative to time-dependence of cytotoxicity. As previously reported, verapamil reduced MeHg-induced cytotoxicity by ~10% at 1 hr of exposure, however, lost its protective effect at 3 hrs (p=0.08). Similarly, nematidine-A (0.25-1 μM) protected neurons against MeHg toxicity at 1 hr (p=0.033), but not 3 hrs of exposure. The reduction of MeHg-induced cytotoxicity by L-type Ca2+ channels contribute to MeHg-mediated neuronal cell death in C. elegans, just as they do in mammalian neurons. The lack of correlation between time and exposure to MeHg may result from upregulation of genes for proteins involved in metal detoxification. Using C. elegans neuronal cultures as a model system for MeHg neurotoxicity may help uncover why certain neurons are more susceptible to MeHg-induced cell death furthering our understanding of the underlying mechanisms.

1457 MTBE Disrupts HIF1-Vegf Regulated Angiogenesis in Zebrafish (Danio rerio).

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Understanding the sensitivity of developing vascular networks to toxic insult is important to advancing vascular biology. Methyl tert-butyl ether (MTBE) induces vascular lesions in the zebrafish embryo, including pooled blood in the common cardinal vein (CCV), cranial hemorrhages (CH), and abnormal intersegmental vessels (ISV). The transcript levels of two isoforms of vascular endothelial growth factor, vegf and vegfc, as well as a primary receptor, vegfr2, are significantly decreased during the critical period (6-somites to Prim-5). The vascular lesions were hypothesized to be a result of the MTBE-induced down-regulation of vegf. An over-expression study was conducted to rescue MTBE-induced vascular lesions. Over-expression of zebrafish vegf resulted in 46% fewer animals exhibiting MTBE-induced CH and 35% fewer embryos exhibiting abnormal ISV, while no rescue was observed for the CCV lesion. Global gene expression changes during the critical period, assayed with the Affymetrix GeneChip® and analyzed with Ingenuity Pathway Analysis®, identified the cardiovascular system as a primary pathway altered by MTBE exposure, as well as other pathways associated with hypoxia inducible factors (HIFs). Two further rescue studies were designed to block HIF1α degradation to test the hypothesis that MTBE toxicity was HIF1α-dependent. Chemical inhibition of HIF1α degradation, by blocking prolyl 4-hydroxylase activity with N-oxalylglycine, rescued both the CCV lesion (24%) and CH (32%). Knockdown of a ubiquitin ligase component, von Hippel-Lindau protein, with an anti-sense morpholino rescued only the CCV lesion (35%). Rescue of MTBE-induced vascular lesions by over-expression of vegfr and inhibition of HIF degradation demonstrated that MTBE toxicity is mediated by a down-regulation of HIF driven Vegf at a critical period during the developing cardiovascular system. Chemicals with anti-angiogenic properties, such as MTBE, can be used to advance the science of angiogenesis in both a disease state and during development. Funding: ES07148, ES05022.
1458 An In Vitro Placental Model Using BeWo Cells for the Prediction of Placental Transfer of Compounds.

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The use of reverse dosimetry for in vitro-in vivo extrapolation (IVIVE) enhances the use of in vitro toxicity data for risk assessment (Louise et al., 2010). With reverse dosimetry, in vitro toxic effect concentrations are translated to in vivo doses using physiologically based kinetic (PBK) modelling. For the PBK models needed for reverse dosimetry, parameter values for kinetic processes need to be obtained, such as for intestinal absorption, metabolism and placental transfer. The present study aims to assess whether an in vitro model for the placental barrier, consisting of human placental BeWo cells, could be used to predict the placental transfer kinetics of compounds. To this end, BeWo cells were grown on transwell inserts to form a confluent monolayer, thereby separating an apical maternal compartment from a basolateral fetal compartment. For a set of 9 compounds the transport velocity from the apical to the basolateral compartment was determined in this model. Relative transport rates obtained in this model were compared with the relative transport rates of these compounds calculated from data of ex vivo human placental perfusion studies reported in the literature (Hewitt et al., 2007). The relative transport rates in the BeWo model were in good correlation (R²=0.95) with the relative transport rates calculated for the ex vivo model (Li et al., submitted). This indicates that the BeWo model can be useful in the prediction of the transplacental transfer of compounds and for obtaining model parameter values for PBK that can be used for reverse dosimetry.

1459 Controlled Hemodynamics and Transport in Primary Hepatocytes Shift Induction and Toxic Responses to Drugs Closer to In Vivo Concentrations.


Preclinical in vitro drug screening systems exhibit efficacy and toxicity responses to concentrations very different from corresponding clinical or in vivo plasma Cmax levels, contributing to poor in vitro-in vivo correlations. We established a primary hepatocyte system using controlled hemodynamics and transport that retains polarized morphology and metabolic function more stably than traditional static cultures. We tested the hypothesis that restoring these critical parameters could achieve in vitro hepatocyte drug response at concentrations closer to in vivo levels. Fresh rat hepatocytes were cultured under controlled hemodynamics alongside static controls for 5 days before treating with various concentrations of test drugs (dexamethasone, acetaminophen, chlorpromazine) for 2 days. Cytotoxic response was evaluated by MTT and ATP assays and cytochrome p450 activity by standard kits. Under controlled hemodynamics, dexamethasone was significantly more toxic to primary hepatocytes at concentrations used for induction studies in static cultures (93.3±5.2%/6 vs. 32.8±2.5% at 50 μM, p <0.01). Induction responses of Cyp3A activity equivalent to 50μM of dexamethasone in static cultures were seen at 2.5 μM under controlled hemodynamics. Cytotoxic dose response curves of acetaminophen demonstrated a leftward shift with IC50 values at 10 μM in devices compared to 30 μM in static cultures. A similar shift in laboratory animals, and the need to extrapolate results from animals to humans. In vitro tissue mimics are a promising avenue for studying the effects of toxicants. Such systems can serve as reliable models in vivo structures and can be systematically probed with a wide range of cues. Since they are engineered, experimentation on these mimics is considerably less complex than those needed to probe tissues and organisms in vivo. We have designed a novel 3D organotypic liver model assembled with three cell types (hepatocytes, sinusoidal endothelial cells and Kupffer cells) and a polyelectrolyte multilayer (PEM) that mimics the Space of Disse. We have established a dose range (20–40μM) for acetaminophen in 3D liver models that is non-lethal yet is capable of perturbing hepatic function. When acetaminophen was administered 4 days after hepatocytes were obtained from rat livers, conventional monolayer, collagen sandwich and 3D liver model cultures showed a small decrease in cell viability. When the drug was administered at day 12, only the 3D liver models showed a decrease in viability. All cultures exhibited a decrease in albumin production and urea secretion. The 3D liver models exhibited better urea secretion than monolayer and collagen sandwich cultures. Intracellular glutathione levels were measured for all three cultures. 3D liver model exhibited approximately 40% decrease in glutathione in comparison to monolayer and collagen sandwich cultures that exhibited greater than 60% decrease. These trends suggest that the inclusion of non-parenchymal cells in the 3D liver model may impart cytoprotective effects and present an in vivo like environment.

1460 Validation of In Vitro Systems to Explore Mechanisms of Striated Muscle Toxicity.

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The validation of in vivo cardiac and skeletal muscle models of toxicity may find a utility in predicting toxicity early within the research and development process. The primary aim of this project was to investigate the extent of translation from an in vivo to an in vitro, system, using a toxicogenomics approach. Female Han Wistar rats were dosed orally via the diet with sulfonyl toxozoline (SI) chemicals for 4 or 28 days. Histopathological evaluation revealed a dose-dependent myositis and myodegeneration in striated muscle. Gene expression profiling of cardiac and skeletal muscle tissues taken from sub-toxic doses identified a number of perturbed cellular pathways including; mitochondrial dysfunction, altered energy metabolism, oxidative stress, cell cycle and apoptosis. This data provided a plausible hypothesis which suggested mitochondrial toxicity as a principle mechanism of SI-induced striated muscle toxicity. To investigate the translation of this toxicity to an appropriate in vitro system cardiac (H9c2) and skeletal muscle (L6) cell lines were selected. A flow based assay was developed in which the cardiac and skeletal muscle cells were adapted to use mitochondrial oxidative phosphorylation rather than glycolysis. This system identified the SI compounds as mitochondrial toxicants. In addition there was a significant increases in mitochondrial reactive oxygen species, which provided further evidence of mitochondrial perturbation. SI treatment also induced cellular hypertrophy; accompanied by cell cycle arrest, and subsequent caspase-mediated apoptosis. These in vitro results were consistent with the transcriptomics data, providing further validation that the cell systems models may have utility for predicting striated muscle toxicity. Future work will be aimed at exploring tissue-specific toxicity using higher tier in vitro models (liver, muscle bionetwork flow through systems) which might better represent the in vivo system.

1461 3D Liver Models for Investigating Drug-Induced Hepatotoxicity.

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Testing the toxic effects of chemicals has traditionally relied on large-scale animal studies. In addition to being extremely expensive, this strategy has come under scrutiny due to an over-reliance on animal models resulting in lower than expected success rates of translating results from animals to humans. In vitro tissue mimics are a promising avenue for studying the effects of toxicants. Such systems can serve as reliable models in vivo structures and can be systematically probed with a wide range of cues. Since they are engineered, experimentation on these mimics is considerably less complex than those needed to probe tissues and organisms in vivo. We have designed a novel 3D organotypic liver model assembled with three cell types (hepatocytes, sinusoidal endothelial cells and Kupffer cells) and a polyelectrolyte multilayer (PEM) that mimics the Space of Disse. We have established a dose range (20–40μM) for acetaminophen in 3D liver models that is non-lethal yet is capable of perturbing hepatic function. When acetaminophen was administered 4 days after hepatocytes were obtained from rat livers, conventional monolayer, collagen sandwich and 3D liver model cultures showed a small decrease in cell viability. When the drug was administered at day 12, only the 3D liver models showed a decrease in viability. All cultures exhibited a decrease in albumin production and urea secretion. The 3D liver models exhibited better urea secretion than monolayer and collagen sandwich cultures. Intracellular glutathione levels were measured for all three cultures. 3D liver model exhibited approximately 40% decrease in glutathione in comparison to monolayer and collagen sandwich cultures that exhibited greater than 60% decrease. These trends suggest that the inclusion of non-parenchymal cells in the 3D liver model may impart cytoprotective effects and present an in vivo like environment.

1462 In Vitro Cell Models: Moving from Tumor Cell ‘Monsters’ to Differentiating Immortalized- or Stem Cells.

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The Tox21 Consortium aims to develop in vitro quantitative High Throughput Screening (qHTS) using human cells and targets to replace animal testing. In screening for cytotoxicity, primary human cells make useful models, but results from primary cells are irreproducible and the cells are problematic to obtain. At the other extreme, transformed cell lines are readily available and reproducible, but are often poor representations of normal human tissues. Between these extremes, immortalized cells or differentiated stem cells provide reliable phenotypic models for neurones, kidney proximal tubule epithelial cells or podocytes, cardiac myocytes, vascular endothelial cells, and hepatocytes. We are evaluating cell models for these cell types by evaluating susceptibilities to toxicants that are organ-selective in vivo. Efforts are also underway to adapt such cells that require differentiation to qHTS in 1536–well plate format. Gene expression profiling is being leveraged to dissect the modes of cytotoxicity, and to determine why in vitro models succeed for some compounds and fail for others. We will present the evaluation of several such toxicological models vis-à-vis: differentiated phenotypes, cytotoxicity assays, and gene expression profiling.
BSEP (Bile Salt Export Pump) inhibition has been proposed as a mechanism for drug-induced cholestasis, a subtype of drug induced liver injury (DILI). Screening systems for BSEP inhibition have been established in membrane vesicles from S9 insect cells over-expressing rat or human BSEP transporter. However, as any cell-free assay, the data from vesicle assays might not reflect true BSEP inhibition profile due to lack of biotransformation and metabolism. In addition, the contribution of MRP2 (Multidrug resistance-associated protein 2) to drug induced liver cholestasis also needs to be considered.

In this study, we examined 18 potential cholestatic compounds, which were either known to be BSEP inhibitors, dual inhibitors for BSEP and MRP2 or glutathione depletors, in both human and rat BSEP and MRP2 vesicle assays. Moreover, a high-content imaging assay using fluorescent selective substrates for BSEP and MRP-2 transporters including CFMDA and CLF, was used to examine these compounds in inhibition of bile canalicular excretion in both human and rat sandwich-cultured hepatocytes. In addition, CFMDA and CLF were characterized for their ability as uptake substrates for BSEP and MRP2 transporters. We found that both, CFMDA and CLF were not selective for BSEP transport since they were also potent MRP2 substrates. Vesicle and hepatocyte canalicular inhibition data showed no substantial species difference between human and rat, for either transporter.

Furthermore the vesicle data correlated well with the hepatocyte canalicular data. However, for a few compounds the vesicle data did not translate into hepatocyte canalicular data probably due to impact of cell-based effects such as metabolism or permeability.

In conclusion, to better predict drug-induced cholestasis, a larger set of compounds with known clinical DILI outcomes need to be tested in both vesicle and hepatocyte canalicular assays in order to elucidate the mechanisms involved in liver toxicity.

**1464 Development of Alternative In Vitro Methods to Screen for Pulmonary Toxicities—Characterization of Epithelial-Macrophage Coculture and In Vitro Assay Conditions at the Air-Liquid Interface (ALI).**

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The successful development of in vitro assays with cultured pulmonary cells and aerosol is instrumental for creating toxicity/screening tests for use during product development. To optimize for future in vitro aerosol exposures using the NanoAerosol Chamber for In Vitro Toxicology system, rat lung epithelial L2 cells and NR8383 macrophages (MO) were co-cultured on an ALI system and cell metabolism (XTT), cytotoxicity (LDH), and cytokine (IL-6) release assays were developed as endpoints to assess cell toxic effects following particle exposures. Co-cultures were exposed to 0-80 μg/cm² ZnO fine particles in supplemented growth medium for 2 hrs followed by 24-hr maintenance of cultures at ALI. Results showed that both XTT and LDH assays performed with cells co-cultured at the ALI were able to detect toxicity in a dose-response manner. Cytotoxic effects measured following 20 μg/cm² ZnO particle exposures (i.e., <1%) increased with epithelial cell/MO (L2:NR8383) ratios, revealing that the seeding ratio strongly affects the assay sensitivity. In a separate study, optimization for cytokine release assay performed with different cell seeding densities and L2:NR8383 ratios showed the highest IL-6 release (~2.5 ng/ml/cm²) from cultures seeded at 0.35/cm² in a 2:1 ratio and maintained at ALI for 1 day. Maximum stimulation of IL-6 release was detected from the optimized cultures 4- and 24-hr post-exposure to low dose (2.5 μg/cm²) ZnO (200-300% of medium-exposed controls); IL-6 release was suppressed to 0-50% of controls after relatively high-dose ZnO (>10 μg/cm²) exposures. In conclusion, these studies are useful in characterizing and optimizing the co-culture and in vitro assay conditions with ALI which could better simulate in vitro exposure of lung cells to inhaled materials (when compared to a submerged exposure system) and ultimately should facilitate the development of reliable in vitro cell exposure systems and cell-based lung toxicology/screening tests.

**1465 Predicting Heart-Specific Toxicity Using Two Cell Models: Human iPSC-Derived Cardiomyocytes and Human Liver Cells (HepaRG).**

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A major reason for the failure of new drugs is due to adverse effects in the cardiovascular system. An in vitro model capable of identifying heart-specific liabilities would be of considerable value. To differentiate heart toxicity from liver toxicity, a dual cell model was developed that utilizes changes in cell health and function following exposure to a test drug. Human cardiomyocytes and normal human liver cells were used as the test system. Cardiomyocytes were derived from induced pluripotent stem cells (iPSC) obtained from Cellular Dynamics International. Terminally differentiated bi-potent HepaRG cells were from Life Technologies. Each cell type was established in a 96-well format and prepared in triplicate wells for each concentration. Markers of cell health (ATP, LDH leakage, and GSH) were monitored in both cell types. Additional information was collected for beat rate (BR) in heart cells and endpoints were monitored over concentration and time. The concentration-response curves were compared using mean IC50 values. Beat rate (BR) was measured using the xCelligence RTCA Cardio system (ACEA). To test the model, hepatoxic (camptothecin (CAMP), rotenone (ROTI)) or cardiotoxic (doxorubicin (DOX), mitoxantrone (MTX)) compounds were used. These were added to cells at concentrations of 0, 0.1, 1, 10, and 100 μM and exposed for 24 hr. In addition, drugs known to produce QT prolongation (Tetraenadine (T) and verapamil (V)) were included and the BR determined. T and V significantly reduced BR at a concentration of 0.3 μM. To determine heart (H) or liver (L) specificity, average IC50 values were obtained for each cell type and the H-to-L ratio calculated. H-to-L ratios were MTX = 0.5, DOX = 2.6, ROT = 0.02 and CAMP undetermined. Ratios < 1.0 indicate cardiotoxicity, those >1.0 but < 3 indicate toxicity in both models, while ratios > 3 indicate hepatotoxicity. BR data was used to improve these predictions. The combined data provided better resolution and enabled cardiac toxicity to be determined with greater confidence.
Phagotest® (Glycotope Biotechnology, Berlin, Germany) is routinely used in clinical toxicity studies to assess phagocytosis in humans. It allows the quantitative determination of leucocyte phagocytosis by measuring the percentage of phagocytes that have ingested bacteria and the number of ingested bacteria per cell. In the present study, 2 ml samples taken from 6 male and 6 female Sprague-Dawley rats onto heparin lithium tubes were incubated with FITC-labeled E.coli bacteria at +37°C. The phagocytosis was stopped by placing the samples on ice, adding a quenching solution. After washing and erythrocyte lysis, a DNA staining solution was added prior to flow cytometry analysis using a XL MCL flow cytometer with the Expo 32 ADC software (Beckman Coulter, France). The percentage of phagocytizing cells (granulocytes and monocytes, differentiated by side scatter and forward scatter) and their mean fluorescence intensity (number of ingested bacteria) were determined in four independent runs. The mean ± SD of the percentage of phagocytizing granulocytes was found to be 84.4 ± 5.0 for males and 86.2 ± 11.02 for females, and the mean ± SD of the percentage of phagocytizing monocytes, 51.6 ± 8.3 for males and 55.5 ± 11.7 for females. Intraclass-variation (CV%) was between 10.4% and 13.6% for males, and between 18.1% and 24.2% for females. These results demonstrate that the Phagotest® kit can be used to assess phagocytosis in rats as it is in humans.

**1468 Thirdhand Smoke: Extraction and Cytotoxicity.**

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Thirdhand smoke (THS) consists of chemicals that remain in indoor environments after secondhand smoke has cleared. The purpose of this study was to develop an extraction protocol for THS and evaluate its cytotoxicity using terytchol exposed to cigarette smoke for 110 hours over 334 days and stored at room temperature (RT). THS was extracted in DMEM culture medium at different temperatures and for different lengths of time, and concentrations of nicotine, and related chemicals in the extracts were measured. Highest concentrations of nicotine, cotinine, nicotilline, anatalline, N-formylnornicotine, 2,3-bipyrindine, N-nitrosornicotinimide (NNN) and 4-(methylisotrosinosimide)-1-(3-pyridyl)-1-butanone (NNK) were present in the extracts made at RT for 1 hr. However, highest concentrations of 1-(N-methyl-N-nitrosamino)-1-(3-pyridyl)-4-butanal (NNYA) and myosmine were in the 1 hr extract made at 4°C. The 1 hr, RT extract had 144 μg of nicotine per gram of terytchol. A terytchol bathrobe, weighing 450 g and similarly exposed to smoke, would contain 64 mg of nicotine, which is higher than the LSD50 dose for humans. Cytotoxicity was assayed using mouse neural stem cells. The extract made at RT for 1 hr was more cytotoxic than that made for 2 hrs. However, when extracted at 4°C, the 2 hr extract was more cytotoxic. Prolonging the extraction time to 24 hours at RT caused complete loss of cytotoxicity, whereas extracting for 24 hours at 4°C helped to preserve some activity. Extracts made with lower headspace volume were more cytotoxic than when the headspace volume was large, presumably due to better preservation of volatile chemicals. In general, the extract made for 1 hr at RT was most cytotoxic, killing cells at the full strength and 30% dose. These data are consistent with the interpretation that cytotoxicity of THS extracts resides in the volatile fraction and that extraction conditions are critical when quantifying chemicals in THS.

**1469 Prevalidation of the Ex Vivo Model Precision Cut Lung Slices (PCLS) for the Prediction of Respiratory Toxicology.**

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Introduction: For acute inhalation toxicity studies, animals inhale chemicals. At the beginning it is difficult to estimate non-toxic doses for *in vivo* inhalation. In the context of REACH and the principle of 3Rs, there is a public demand for alternative methods. The goal of this BMBF-funded project is the pre-validation of PCLS as a suitable *ex vivo* alternative model to replace pre-studies of inhalation toxicity. The project is conducted in three independent laboratories. BfR provides support in biostatistics.

Methods: In all participating laboratories, PCLS were prepared and exposed to 5 concentrations of industrial chemicals in DMEM under standard condition for 1 hour. After post-incubation, chemical-induced toxicity was assessed by LDH and WST-1 assay. In addition, PCLS protein content and pro-inflammatory cytokine IL-1β were measured by BCA assay and ELISA, respectively. For all endpoints a sigmoid dose-response model was fitted to the data and EC50 values were calculated. Test acceptance criteria were established for each endpoint.

**1470 High-Content Analysis of Drug-Specific Neurotoxic Effects on Rat Dorsal Root Ganglion Cells.**

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The incidence of chemotheraphy-induced peripheral neuropathy (CIPN) is increasing because more neurotoxic drugs are being developed and these are increasingly administered in combinatorial regimens to longer living cancer patients. To evaluate neurotoxic effects, a high content analysis (HCA) strategy was developed to assess response differences of exposed rat dorsal root ganglion cells (DRG). DRG microwells were exposed to CIPN-inducing drugs (up to 1 μM) for 24 hr. After exposure, treatment medium was removed and cells were fixed, or were given fresh culture medium and allowed to “recover” for 24 or 72 hr prior to fixation. DRG were fluorescently stained for nuclei, acetyl-β tubulin, and Nissl bodies. Four of the six drugs evaluated caused concentration-specific changes in acetyl-β tubulin staining (AATS) and total neurite area (TNA) after 24 hr exposure. Eribulin and colchicine were most potent, reducing TNA by 50% at <10 nM, but only colchicine effects were reversible after drug removal (AATS intensity doubled in vivo experiments in three independent laboratories, concentration dependent toxicity could be shown for aniline, guanidinium, triton X-100 and paracetamol, but not for lacosate and methyl methacrylate. EC50 values obtained for the WST-1, LDH and BCA data were very similar in all participating laboratories. No increase in IL-1β was observed for these chemicals.

Conclusion: Local respiratory toxicity induced by chemicals could be tested with comparable results in the PCLS model without *in vivo* experiments in three independent laboratories. The standardization of the PCLS method was successful and the reproducibility of the results is very promising after testing of the first 6 substances.
1472 A Transcriptomic Comparison between the Neural and Cardiac Embryonic Stem Cell Tests (ESTn and ESTc).

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In vitro screening assays may increase testing efficiency and reduce animal use in developmental toxicity testing. The cardiac mouse embryonic stem cell test (ESTc) is a promising in vitro assay in this field, in which the effect of developmental toxicants on cardiomyocyte differentiation is assessed. To increase prediction of the EST approach, we developed a neural differentiation variant of the stem cell test (ESTn). In both ESTn and ESTc, we performed a series of transcriptomic studies to characterize gene expression changes 1) across time during normal differentiation and 2) in response to a series of developmental toxicants in the ESTn and ESTc. In the present study, gene expression profiles of ESTn and ESTc over time as well as model-specific changes induced by seven compounds are compared. Time-related gene expression profiles showed that specific genes changed over time differently in each model, related to the two specific lineages of differentiation. Interestingly, compound-induced gene-expression changes were generally model-specific, particularly for methylmercury and fluazinamide, which were predicted better in ESTn and in ESTc, respectively. Valproic acid-induced gene expression changes were most comparable between ESTn and ESTc. Direct transcriptomic comparisons between the ESTn and ESTc models indicate that both assays support and complement each other. Therefore, a combined transcriptomics approach, incorporating ESTc and ESTn, may result in improved developmental toxicant detection over individual assays.

1473 In Vitro Proliferation in Human and Rat Urothelial Cells.

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Despite the regular use of the rat in toxicity studies, which are used for human risk assessment, the extrapolation from rat to humans is often difficult. This study set out to explore the utility of cell culture systems to better understand species differences. As urine is a main route of chemical excretion from the body, the epithelial barrier lining of the bladder (urothelium) may be exposed to higher concentrations of xenobiotics, with the potential for adverse findings. Methods have been successfully developed to culture normal human urothelial (NHU) cells in vitro, with subsequent differentiation to form a functional barrier. With this in mind, equivalent normal rat urothelial cell culture systems were grown from Wistar (NRU-W) and Homozygous Scottish (NRU-HS) rats. They were compared to human cells culture by exploring their regulation and capacity for proliferation. NHU cells could be propagated beyond passage 6 in serum-free medium, while NRU-W could only be maintained up to passage 4, and NRU-HS cells required serum to grow to passage 1. By immunofluorescence, NHU cells were shown to express nuclear Ki67 and cyclin D1, but the same proliferation markers were cytoplasmic in NRU cells. In contrast, Ki67 was virtually negative in histological sections of human and rat urothelium, but strongly expressed (nuclear) in Wistar. The results for human urothelium reflect the fact it is mitotically-quiescent, and has a low rate of turnover in vivo, but has a large capacity for regeneration. As urine is a main route of chemical excretion from the body, the epithelial barrier lining of the bladder (urothelium) may be exposed to higher concentrations of xenobiotics, with the potential for adverse findings. Methods have been successfully developed to culture normal human urothelial (NHU) cells in vitro, with subsequent differentiation to form a functional barrier. With this in mind, equivalent normal rat urothelial cell culture systems were grown from Wistar (NRU-W) and Homozygous Scottish (NRU-HS) rats. They were compared to human cells culture by exploring their regulation and capacity for proliferation. NHU cells could be propagated beyond passage 6 in serum-free medium, while NRU-W could only be maintained up to passage 4, and NRU-HS cells required serum to grow to passage 1. By immunofluorescence, NHU cells were shown to express nuclear Ki67 and cyclin D1, but the same proliferation markers were cytoplasmic in NRU cells. In contrast, Ki67 was virtually negative in histological sections of human and rat urothelium, but strongly expressed (nuclear) in Wistar. The results for human urothelium reflect the fact it is mitotically-quiescent, and has a low rate of turnover in vivo, but has a large capacity for regeneration, which can be seen in culture. It appears that there may be fundamental differences in the way in which rat urothelium is regulated, with cells maintained in the cell cycle in vivo and with a less proliferative phenotype observed in vitro. Future work will aim to further characterise these differences, and to understand the consequence of these findings for the development of a comparative model.

1474 In Vivo-In Vitro Comparison of Respiratory Tract Toxicity Using Human 3D Airway Models and Human A549 and Mouse 3T3 Monolayer Cell Systems.

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Four in vitro systems to predict acute inhalation toxicity were evaluated. 19 substances (lactose, paracetamol, methylmethacrylate, aniline, acetic- and trimellitic anhydride, N-ethylchloroformate, octanol chloride, isophorone and toluene diisocyanate, zinc oxide, paraquat, glutar- and formaldehyde, acetone, ethanol, diethylformamide, ammonium hexachloroplatinate and sodiumdodecylsulfate) were tested in 3D human airway epithelial models, EpiAirwayTM and MucilAirTM, and in A549 and 3T3 monolayer cell cultures. Cytoxicity was assessed by determining LDH release and MTT or WST reduction. IC50 values were compared to literature rat 4-hour LC50 values classified according to the US EPA and GHS hazard categories. Best results were achieved with a prediction model identifying non-toxic substances (determination of LDH release in 3T3 cells: sensitivity 1.00 and specificity 0.89).

Further predictions of in vivo hazard categories based on four in vitro hazard categories resulted in mediocre correlations: 9 of 19 test substances were classified concordantly in the MucilAir™ system determining MTT reduction and 8 of 19 in A549 cells determining WST reduction. Concordance could be improved by excluding substances leading to pulmonary edema and emphysema in vivo. None of the test systems was outstanding and there was no evidence that the use of 3D or monolayer systems using respiratory tract cells provide an added value to simple 3T3 monolayers. However, the test systems only reflected bronchiol epithelia and alveolar cells and only investigated cytoxicity so that effects occurring in other cells by other mechanisms were not recognized.

1475 Neurotoxicity In Vitro: Assessment of the Predictivity of Neuronal Networks Coped to Microelectrode Arrays for Identification of Neurotoxins.

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A challenging aspect to assure the safety of a product is the assessment of its neurotoxic hazard potential. Currently, only in vivo methods are regulatorily accepted and so far, no in vitro model has been fully validated. The majority of the test systems are reduced to the analysis of cytotoxicity in immortalized cell lines, without including unique characteristics of the nervous system, such as axonal transport, synaptic transmission or its electrophysiology. Recently, with the advance in technology and the ability to maintain neuronal models for prolonged periods, a test system emerged, combining the use of microelectrode arrays (MEAs) and in vitro culture of 2D neuronal networks (NN). Herein, we report on the in-house validation of the NN MEA assay using a set of 43 compounds with known neurotoxic and non-neurotoxic potential with the aim to use it for screening of compounds under development. The results demonstrate that the methods present a sensitivity of 71%, while the specificity is still an aspect for optimization, since in its current status, it cannot distinguish specific neurotoxicity from unspecific cytotoxicity. In order to increase the sensitivity and predictivity, we are currently working on the combination of the electrophysiological assessment with a panel of cytotoxicity assays.

1476 A Human 3D Myocardial Microtissue Model for Cardiotoxicity Testing.

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Cardiomyocytes (CMs) are terminally differentiated cells in the adult organism and regeneration is limited. This is a worrisome fact, since insults such as ischemia and cardiotoxic compounds can lead to cell death and irreversible reduction of cardiac function. As testing platform, isolated organs and primary cells from rodents have been the standard in research and toxicology so far, but due to a very limited cell supply there is a strong need for better in vitro models. Hence, an in vitro model comprising the advantages of 3D cell culture and the availability of induced pluripotent stem cells (iPSC) from human origin was developed and characterized. Myocardial microtissues (MTs) were generated by self-assembly in multi-well hanging drop cultures. iPSC-derived CMs were evaluated regarding cardiac features and toxicological response. Prior use of the iPSC-derived cells, CMs were characterized after 10 days in standard culture which showed highly differentiated myofibers positive for sarcomeric proteins such as myomesin, myosin, cardiac actin and desmin. The cells showed spontaneous contractile activity and connexin-43 positive gap junctions. In the hanging drop cultures, iPSC-derived CMs formed MTs within 4 days, contracting up to 3 weeks recorded by optical motion tracking. Excited by electrical field pacing they compiled to the external stimulus up to 2Hz. Effects of a cardiac cancer therapeutics such doxorubicin on iPSC-derived CMs in 2D- and 3D-culture were further evaluated with respect to caspase activation, LDH release and cellular ATP levels. Adult human CMs are a very rare and hard to handle cell source. New concepts have to be developed to create cardiac models systems to be used for cardiox testing, ideally in a standard multi-well format. Within this "proof-of-concept" study a novel 3-dimensional human myocardial MT model with developed and characterized iPSC-derived CMs. Morphological and functional characterization underline that this model might become a valuable tool for substance safety testing in the future.
1477 Use of An In Vitro Flow Cytometric Method As a Replacement for Live Animal Experimentation.
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Scope:
Animal welfare guidelines place more and more emphasis on the 3 Rs principle as a way to achieve excellence in animal care and use. In that objective, we have developed an in vitro flow cytometry method that allows us to quantify the immune response to IL-2 stimulus in a lymphocyte population replacing the use of live monkeys.

We have evaluated the potency of test items to phosphorylate Stat5 in lymphocyte subsets expressing or not the IL-2 high affinity receptor subunit (CD25).

Experimental Procedures and Results:
Cytomolgous monkey peripheral blood mononuclear cells (PBMC) were stimulated with increasing doses of the test items under evaluation. The PBMCs were then stained using conjugated monoclonal antibodies specific to different lineage markers such as CD3, CD4, CD8 and CD25 to allow distinction between the different T lymphocyte subsets. In addition, in order to evaluate the potency of the test items to differentially stimulate CD25+ and CD25- lymphocytes, we have evaluated the activation of Stat5 by means of the measurement of the phosphorylation of the Signal Transducer and Activator of Transcription (Stat5).

Conclusions:
This innovative approach enables us to replace animal experimentation by an in vitro assay. Our method assesses the potency of test items to signal through the IL-2 receptor in different lymphocyte subsets.

1478 The Role of Facilitated Transport by Serum Protein in In Vitro Intrinsic Clearence.

When a chemical is exposed to an in vitro cell assay in culture medium with serum protein, the effect (e.g. clearance) observed may be lower than when the chemical is exposed in culture medium without serum protein. This is because the chemical can bind to serum protein, which reduces the unbound free concentration of the chemical available for uptake into cells. Therefore, it is better to determine in vitro intrinsic clearance (CLint) based on unbound fractions of chemicals. However, a few studies have suggested that serum protein may also facilitate the transport of chemicals through aqueous media (facilitated transport). Thus the presence of serum protein may increase the uptake rate of chemicals into cells and solid phase microextraction (SPME) fibers. If the uptake rate determines clearance, then the presence of serum protein may increase clearance, thus hampering the extrapolation of in vitro CLint to in vivo clearance when clearance assays use varying concentrations of serum. Therefore, the uptake rate and clearance of the strongly-albumin bound, quickly cleared chemical tylosin was measured in HepaRG, HepG2 and H4IIIE hepatoma cell lines at varying concentrations of bovine serum albumin using the substrate depletion approach. To measure the free fraction, mimic uptake of tylosin into cells, and facilitate the modeling of the uptake into cells, a SPME method was developed for extracting the unbound chemical from the exposure medium. Results indicate that in vitro CLint of tylosin increases with increasing albumin concentration when using the unbound fraction. SPME was successfully applied to determine the free concentration and study the uptake rate of unbound tylosin into cells.

1479 Screening of Cosmetics Ingredients for Phototoxic Potential Using the In Vitro 3T3 Neutral Red Uptake Phototoxicity Test.
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Phototoxicity is an acute toxic response after exposure to a phototoxiant and either UV radiation or visible light (UV/VIS). Phototoxicity from substances applied topically typically occurs at the site of photo-irradiation. Phototoxicity is the result of direct cellular damage caused by a nonimmunological inflammatory response. Clinical symptoms of phototoxicity resemble an aggravated sunburn (erythema, increased skin temperature, pruritis and edema). Phototoxicity reactions have been reported for both synthetic substances and those which occur naturally (e.g., botanical extracts). Although symptoms generally subside quickly, the potential for substances used in topical products to cause phototoxicity is clearly of concern for manufacturers of cosmetics, personal care and other consumer products. Indeed, the potential to cause phototoxicity from substances applied topically was evaluated by utilizing various animal models; however in 1997 the 3T3 Neutral Red Uptake Phototoxicity Test (3T3 NRU PT) was validated by ECVM’s Scientific Advisory Committee as an in vitro method for evaluating the phototoxic potential of chemicals shown to absorb in the UV/VIS range. To illustrate the utility of the 3T3 NRU PT as a useful screening tool in the safety evaluation of potential cosmetic ingredients, the results of the evaluation of 42 botanical extracts and 25 synthetic chemicals found to absorb in the UV/VIS range are reported. Most substances evaluated were found not to be phototoxic in vitro; however, 9 substances were identified as potentially/probably phototoxic in the 3T3 NRU PT and were eliminated from further consideration for use as cosmetic ingredients. Several substances found to be non-phototoxic in the 3T3 NRU PT were formulated with other ingredients in a prototype cosmetic formulation and subject to clinical testing. No manifestations of phototoxicity were observed in any of the test subjects in the prototype formulation containing any of the substances identified as non-phototoxic in vitro.

1480 Validation of a Cell Microelectronic Sensing Technique for Semi-Quantification of Cyanobacterial Toxins (Microcystins) in Recreational Water.
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Microcystins (MCYSTs) are hepatotoxins produced by cyanobacteria (blue-green algae) commonly found in fresh water. There are a few semi-quantitative and quantitative detection methods available including protein phosphate inhibition assay (PPI), enzyme-linked immunosorbent assay (ELISA) and liquid chromatography tandem mass spectrometry (LC/MS/MS). Our laboratory recently developed a novel assay using a cell microelectronic sensing technology known as RTCA (Real-Time Cell Analyzer, Roche xCELLigence system) for detecting MCYSTs cytotoxicity. The assay is based on the fact that MCYSTs toxicity requires the active uptake of MCYSTs into the cytoplasmic membrane which is mediated by the organic anion transporting polypeptides (OATPs). By comparing the 48-hour cytotoxic effects of MCYSTs in wild type Chinese hamster ovarian cells (CHO/WT) and in CHO cells with OATP1B3 expression (CHO/OATP1B3), we semi-quantified MCYSTs cytotoxicity in recreational water using standard curves prepared from MCYST-LR, the most common MCYST analogue. MCYST levels in water samples were compared with results from PPI, ELISA and LC/MS/MS. The RTCA directly demonstrated MCYSTs total toxicity and the semi-quantitative results have high correlation to those from other methods (p-value < 0.0001), particularly at the Canadian recreational water guideline level (20 μg/L) or greater.

In summary, we have validated a semi-quantitative assay for cytotoxic testing of MCYSTs in recreational water by observing real-time toxic response at microgram per litre concentrations.

1481 Human Intestinal Microflora Reduces Butyl Paraben Toxicity in HepG2 Cell Cultures.
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Parabens are alkyl esters of p-hydroxybenzoic acid, including methyl paraben, ethyl paraben, propyl paraben, and butyl paraben. In the present study, possible role of metabolism by fecalase in BP-induced cytotoxicity was investigated in HepG2 cell cultures. As an intestinal bacterial metabolic system for butyl paraben, human fecal preparation containing intestinal microflora (fecalase) was employed. Initially, among the parabens tested, cytotoxicity of butyl paraben was most severe. When butyl paraben was incubated with fecalase, it rapidly disappeared, in association with reduced cytotoxicity in HepG2 cells. In addition, butyl paraben-incubated with fecalase reduced cytotoxicity of HepG2 cells in a concentration-dependent manner. Moreover, butyl paraben-incubated with fecalase significantly caused an increase in Bcl-2 expression together with a decrease in Bax expression and cleaved Caspase-3. Furthermore, anti-apoptotic effect by the incubation of butyl paraben
with fecalase was also confirmed by the terminal deoxynucleotidyltransferase-mediated dUTP-biotin nick-end labeling assay. Taken all together, the findings suggested that metabolism of butyl paraben by human fecalase might have protective effects against butyl paraben-induced toxicity in HepG2 cells.

1482 Characterization of Cellular and Molecular Markers of Toxicity in C6-Glioma Cells Exposed to Modern Pyrethroids.
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Pyrethroids (Pyr) are insecticides increasingly used in various pest control applications. Low levels of Pyr residues have been detected in diverse environmental samples, food and human urine. In vitro systems may be useful to characterize the diversity of primary and secondary toxicogenetic pathways that may account for the pyrethroid-type specific divergence in clinical syndromes observed in small rodents. We use C6-glioma cell cultures to identify toxicologically relevant Pyr exposures that may require cumulative risk assessment efforts in animals and in vivo. Four compounds (bifenthrin, tefluthrin, α-cypermethrin, deltamethrin) were examined after 24 h of Pyr exposure. We also evaluated pyrethroid actions in C6 cells treated with sodium butyrate (NaBu) which induce an astrocyte-like phenotype. Threshold dose and EC15(24h) estimates were obtained after modeling MTT test data. In order to study subcellular integrity and signaling pathways with greater specificity and sensitivity, Mitotracker-Red and Neutral Red staining (informing on mitochondrial and lysosomal status, respectively), caspases and p53 (cell death related markers), AchE (an insecticide neurotoxicity related target) and GFAP (glia-specific xenobiotic receptor) were then examined. Low μM exposures induced dose-related declines in cell viability in both cell systems. We further found dose-related increases in fragmented mitochondria and mitochondrial membrane potential disruption in C6 cells. Increases in cell death related proteins and drop of GFAP levels were also observed in NaBu-treated cells. In addition, cholinesterase inhibition was not evident below EC15 levels for general cell viability for any test chemical. These results confirm that μM exposures affect cell viability markers unrelated to the proposed primary mode of action of pyrethroids, and that the classically proposed target cell for pyrethroids, the neuron, is not the only cell type that may be susceptible of citotoxicity and severe cell damage in mammals.

1483 DNA Damage-Induced by Goldenseal Constituents.
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Goldenseal is used for the treatment of gastrointestinal disturbances, urinary disorders and inflammation. The major alkaloid constituents in goldenseal are berberine, hydralazine, and canadine. Because it is one of the most widely used herbal dietary supplements in the United States and the lack of carcinogenicity data, goldenseal was nominated to the National Toxicology Program (NTP). The NTP conducted 2 year bioassay on goldenseal and reported that goldenseal increased the incidence of liver tumors in rodents. However, the mechanisms of goldenseal-associated liver carcinogenesis are unclear. In this study, we compared genotoxicity of five goldenseal constituents and studied the underlying mechanisms. Five goldenseal constituents tested (berberine, palmatine, hydralazine, hydralizinine, and canadine) did not induce mutagenicity in the salmonella mutation assay. Berberine and palmatine at high concentrations showed positive results in the mouse lymphoma assay. Berberine and palmatine also caused DNA strand breaks in cultured hepatic cells whereas the rest three did not. The expression of the γ-H2AX, a biomarker of double strand breaks, was induced in a dose-dependent manner in response to berberine treatment. In addition, berberine and palmatine suppressed that activities of both topoisomerase I and II, indicating that the inhibitory effect may contribute to berberine- and goldenseal-induced genotoxicity and tumorigenicity.

1484 DNA Adduct Formation and Mutation Induction by Aristolochic Acid in Rat Spleen.
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Aristolochic acid (AA) is a potent human nephrotoxin and carcinogen. We previously reported that AA treatment resulted in DNA damage and mutation in the kidney and liver of rats. In the present study, we have determined the DNA adducts and mutations induced by AA in rat spleen. Big Blue transgenic rats were gavaged with 0, 0.1, 1.0 and 10.0 mg AA/kg body weight 5-times/week for 3 months. Three DNA adducts, 7-(deoxyadenosin-N6-yl)-aristolactam I and II and 7-(deoxyguanosin-N2-yl)-aristolactam I and II, were identified by P32-postlabeling. Over the dose range studied, there were strong linear dose-responses for AA-DNA adduct formation in the treated rat spleens, ranging from 4.6 to 217.6 adducts/10^8 nucleotides. Spleen cell mutant frequencies (MFs) also increased in a dose-dependent manner from 32.7 to 286.2 x 10^-6 in the treated animals. Mutants isolated from different treatment groups were sequenced; there were significant differences between the spectra of 1 mg/kg AA-treated and control groups, and between the 10 mg/kg AA-treated and control groups. ACT→TA transition was the major type of mutation in AA-treated rats, while GCT→AT transition was the major type of mutation in the vehicle controls. These results indicate that AA is genotoxic in the spleen of rats exposed under conditions that result in DNA adduct formation and mutation induction in kidney and liver.

1485 In Vitro Genotoxicity of Ginkgo biloba Extract and Two of Its Major Components, Quercetin and Kaempferol.

Ginkgo biloba (ginkgo), one of the world’s oldest living tree species, has been used for many years for a variety of medicinal purposes. The NTP 2-year bioassays on ginkgo extract found an increased incidence of liver cancer in mice and thyroid gland cancer in both mice and rats. In this study, the mouse lymphoma assay (MLA) was used to evaluate the in vitro mutagenicity of ginkgo extract and two of its major constituents, quercetin and kaempferol. L5178Y/Tk−/−→3.72 c mouse lymphoma cells were treated with different concentrations of ginkgo extract (0.2-1.2 mg/ml), quercetin (3-30 μg/ml) and kaempferol (2.7-5.7 μg/ml) in the absence of metabolic activation. Ginkgo extract, quercetin, and kaempferol significantly increased the mutant frequency in the MLA. Loss of heterozygosity (LOH) analysis for mutants induced by quercetin and kaempferol were also examined at five microsatellite loci spanning the entire mouse chromosome 11. The results indicated that the mutational spectrum from the quercetin and kaempferol treatment were significantly different from that of the negative control. In addition, the neutral comet assay conducted in the mouse lymphoma cells with quercetin and kaempferol also demonstrated a dose-dependent increase in the DNA double-strand breaks (DSBs). Western blot analysis showed that quercetin increased the phosphorylation of ATM, and consequently increased expression of γ-H2AX, phosphorylated Chk1 and Chk2 in the cells; while kaempferol increased expression of γ-H2AX and phosphorylated Chk1. These results suggest that ginkgo extract and two of its major constituents, quercetin and kaempferol, are genotoxic in the mouse lymphoma cells.

1486 Loss of Heterozygosity Analysis of the Tk Mutants Induced by Cigarette Smoke Condensates in Mouse Lymphoma Cells.
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Cigarette smoke condensate (CSC) is a set of sticky particles comprised of thousands of chemicals created by burning cigarette. Our previous study on 11 cigarette smoke condensates (CSCs) using the mouse lymphoma assay (MLA) has demonstrated that all these CSCs resulted in a dose-dependent increase of both cytotoxicity and mutagenicity in the mouse lymphoma cells. To elucidate the underlying mutagenic mechanism, we examined the mutational types of CSC-induced Tk mutants by analyzing loss of heterozygosity (LOH) using the allele-specific PCR at five microsatellite loci (Tk1, D11Mit42, D11Mit36, D11Mit20 and D11Mit74) spanning the entire chromosome 11. For each CSC treatment, 48 large and 48 small microsatellite loci (Tk1, D11Mit42, D11Mit36, D11Mit20 and D11Mit74) were sequenced; there were significant differences between the spectra of 1 mg/kg AA-treated and control samples. ACT→TA transition was the major type of mutation in AA-treated rats, while GCT→AT transition was the major type of mutation in the vehicle controls. These results indicate that AA is genotoxic in the spleen of rats exposed under conditions that result in DNA adduct formation and mutation induction in kidney and liver.
Cyproterone acetate (CPA), a hormone therapy drug with anti-androgenic activity, is commonly used for androgenisation symptoms in women and treatment of prostate cancer in men. CPA is known to produce liver tumour in rats, with a higher incidence in females. In standard genotoxicity assays, such as Ames test, HGPRT assay, chromosomal aberration assay and in vivo micronucleus assay, negative responses were observed. However, more recent studies have demonstrated that CPA induces a sex-specific genotoxicity, forming DNA adducts in female rats but not in male rats. To investigate the genotoxicity of CPA, and to answer the question if it is sex-specific, we evaluated the genotoxicity of CPA through in vivo Comet assay in both sexes of rats. hOGG1- and Endo III-modified in vivo Comet assay were performed to measure CPA induced oxidative DNA damage. Groups of 5 seven-week old male F344 rats were treated with olive oil or 10, 25, 50 or 100 mg/kg bw CPA in olive oil at 0, 24, and 45 hr. Animals were sacrificed at 48 hr, 3 hr after the last treatment. Liver, testis, kidney and blood were collected for Comet assay. CPA treatment resulted in an increase in DNA strand breaks in the liver of males with significant (p<0.05) increases detected in all the dose groups, suggesting a possible role of reactive oxygen species for CPA-induced genotoxicity. To evaluate the genotoxicity of CPA in female rats, groups of 5 seven-week old female F344 rats will be treated identically with CPA. Liver, mammary gland, uterus, ovary, kidney and blood will be collected for Comet assay and the responses will be compared with the males. This study will provide a further understanding of the cancer mode of action of CPA and if there is any sex-specific genotoxic response in rats.

**1487 Evaluation of Genotoxicity of Cyproterone Acetate in Both Sexes of Rats.**

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**1488 Assessment of Dosimetry and Biological Responses In Vitro Using a Vitrocell® Smoke Exposure System.**

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Routine toxicological assessment of tobacco smoke commonly uses the particulate fraction of the smoke aerosol. The particulate phase of cigarette smoke makes up a small percentage of the total aerosol, approximately 5-9% by weight. The remaining ~91% is associated with the vapour phase of cigarette smoke and is not routinely evaluated. Whole smoke exposure systems are capable of capturing the full interactions of both the particulate and vapour phase together and offer unique potential for toxicological assessment.

In this study we used a Vitrocell® VC10 smoking robot to expose cell cultures at the air-liquid interface to ISO mainstream 3R4F cigarette smoke. All experiments were independent and completed a minimum of three times. For biological assessment we observed a mean fold increase of 5.9 at the highest concentration of smoke deposited mass. QCMs have also acted a important QC tool for smoke exposure experiments and provide valuable information on the exposure system itself.

**1489 Evaluation of the Results of Several Genotoxicity Tests in Carvacrol and Thymol.**

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Carvacrol (CAS499-75-2) and thymol (89-83-8) are fragrance ingredients that have been studied in bacterial and mammalian genotoxicity assays in vitro and in an in vivo chromosome aberration test. These materials have antibacterial and antifungal properties and are cytotoxic. Their cytotoxicity needs to be considered when evaluating the results of genotoxicity assays using mammalian cell cultures because cytotoxicity is the most important confounding factor in analyzing these results. To avoid the occurrence of secondary (indirect) modes of clastogenic activity, the highest concentrations tested in these assays should lead to sufficient levels of cytotoxicity within the guidelines, but not cause extreme cytotoxicity. In published literature, there are inconsistent results of the genotoxicity tests on these two materials, where positive results correlate with cytotoxicity, indicating that positive results can be explained by secondary effects. An SOS Chromotest was least prone to false positives than the Ames test and was negative with both carvacrol and thymol. A positive chromosomal aberration test in rats on both carvacrol and thymol was further evaluated to clarify the cause of the chromosomal aberration observed. Thus an in vitro micronucleus test in human peripheral blood lymphocytes was conducted with careful selection of doses based on cytotoxicity to determine if these materials were potential clastogens or a spindle poison. Both assays yielded negative results, supporting the hypothesis that the previous positive genotoxic outcomes were the result of high cytotoxic activity in the test system and that carvacrol and thymol are likely not genotoxic agents.

**1490 No Direct Genotoxic Potential of Different Radiofrequency Electromagnetic Fields in MRC-5 Cells and B6C3F1 Mice.**

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Public exposure to electromagnetic fields in the radiofrequency spectrum (RF-EMF) has increased dramatically, attracting notice to health risk evaluation. RF-EMF was classified as possibly carcinogenic to humans (Group 2B) by the IARC in 2011. The risk assessment is still limited by data gaps and contradictory data, not only for mobile phones but also for wireless network devices. As there are some indications for disruption of DNA-integrity by RF-EMF exposure, we evaluated the in vitro and in vivo genotoxic potential of different RF-EMF signals by using well established and standardized genotoxicity.

In vitro, normal human MRC-5 lung fibroblasts were exposed intermittently (5 min ON, 10 min OFF) in a 1950 MHz exposure system for 1, 4, and 24 h to various RF-EMF signals (continuous wave (CW), UMTS, WiFi, GSM-basic, RFID) at specific absorption rates (SAR) up to 4.92 W/kg. DNA-strand break (SB) induction and oxidative DNA-damage were subsequently evaluated in an enzyme (hOGG1)-modified comet assay. In vivo, male B6C3F1 mice were whole-body exposed to CW, UMTS, WiFi, or RFID signals at SAR of 1.6, 4.0 and 10 W/kg for 14 d, 20 h/d (5 min ON, 10 min OFF) using a reverberation chamber. Micronucleus (MN) frequency was determined in peripheral blood, immature bone marrow erythrocytes and keratinocytes. Irrespective of signal type, SAR-value, and exposure duration RF-EMF did not mediate significant cytotoxicity, induction of DNA-strand breaks, or oxidative DNA-damage in MRC-5 cells, compared to concurrently sham-exposed cells. In addition, no increase in MN formation was observed in B6C3F1 mice. In conclusion, none of the tested signal modulations revealed evidence for a direct DNA-damaging or cytotoxic potential of RF-EMF.

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**1491 In Vitro Evaluation of Cytotoxic and Genotoxic Effects of Ofloxacin.**

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Ofloxacin (OFX), the second-generation of quinolones, is a broad-spectrum fluorouracil antibiotic used in the treatment of various bacterial infections. In this study, the genotoxic and cytotoxic effects of OFX in cultured human peripheral blood lymphocytes were investigated by measuring chromosome aberrations (CAs), sister chromatid exchange (SCE) assay, mitotic index (MI) and proliferation index (PI). Cultures were treated with three doses of OFX (30, 60 and 120 μg/ml) at 48 h exposure period for all assays. An untreated culture (negative control) and a positive control (positive control) were established, as well. Although the frequency of SCE slightly increased in a concentration-dependent manner, dose-dependent increases were not observed in the CAs. But, these increases were statistically not significant, compared to negative control (P < 0.05). With respect to the cytotoxic effects of the test compound on lymphocyte cultures, as measured by MI and PI, a concentration-dependent reduction in cell proliferation was also observed. The highest dose of OFX (120 μg/ml) significantly reduced the MI values compared to negative control (P < 0.05), whereas decreases in PI values were not found significant compared to negative control.
The results of this study indicate that OFX can exert a cytotoxic effect especially at the higher doses because of statistically significant decreases in MI, but has no genotoxic activity in both CA and SCE assays in human peripheral blood lymphocyte cultures.

Ecatepec County is considered one of the most contaminated counties in the Metropolitan Area of Mexico City. There are reports about the high levels of particulate matter (PM) in this area related to the high traffic and industrial activity. PM is associated with several health problems including genotoxic effects. Children are considered one of the most susceptible populations to air pollution. The aim of this study was to evaluate the DNA damage related to the exposure to PM in the air. We conducted an air monitoring study and an epidemiological survey in schoolchildren from 6 to 10 years (n=86) of both genders in an area of Ecatepec County and the following was evaluated: concentrations of PM10 and PM2.5 in the air, as well as the organic and elemental carbon content (CO and CE, respectively), the presence of microcystine (MN), apoptotic cells (AC) and necrotic cells (NC), and the nuclear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells. The geometric average age was 8.4 years; 41% of children showed at least 1 MN (range 1-6), 27% showed 1-49 AC and 16% had 1-9 clear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells. The geometric average age was 8.4 years; 41% of children showed at least 1 MN (range 1-6), 27% showed 1-49 AC and 16% had 1-9 clear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells. The geometric average age was 8.4 years; 41% of children showed at least 1 MN (range 1-6), 27% showed 1-49 AC and 16% had 1-9 clear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells. The geometric average age was 8.4 years; 41% of children showed at least 1 MN (range 1-6), 27% showed 1-49 AC and 16% had 1-9 clear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells.

Obesity is a global epidemic. Worldwide 500 million people are classified as obese with a global prevalence that continues to rise. Obesity and its co-morbidities are amongst the leading causes of global mortality and morbidity and pose substantial socioeconomic burdens on health services worldwide. Bariatric surgery is a form of gastrointestinal surgery that leads to sustained weight loss, resolution of type 2 diabetes and a decrease in cancer risk. These operations alter gut microbial composition which is reflected in the bacterial composition of faeces. We hypothesise that the protective effect of bariatric surgery is due to a decrease in diet-derived intestinal toxic burden. Here, we determined the effect of bariatric surgery on intestinal cytotoxicity and genotoxicity in man. Aquous dimethyl sulfoxide faecal extracts were obtained from preoperative and 2 month postoperative samples from 5 individuals undergoing Roux-en-Y gastric bypass. Human MCL5 cells, a lymphoblastoid cell line expressing CYP1A1, 1A2, 2E1 and 3A4 and epoxide hydrolase, were used to assess faecal extract toxicity and genotoxicity was determined at the TK and HPRT loci using benzo(a)pyrene as a positive control. We found a trend of decreased cytotoxicity postoperatively, as assessed by relative total growth (20% decrease compared to preoperative faecal extracts). Furthermore, the genotoxicity of the faecal extracts decreased postoperatively at both the TK and HPRT loci (50% and 70% respectively, whereas the BaP control induced mutation frequency more than 50 fold). These results support the hypothesis that bariatric surgery leads to a decrease in diet-derived intestinal toxicity burden, which in turn may contribute to the health beneficial effects associated with surgery.

The metabolism of the tobacco-specific carcinogen NNK form DNA adducts in animal models. One report indicates that NNK could cause damage to the mitochondrial as well as nuclear genome in rats (Stepanov and Hecht, 2009 Chem. Res. Toxicol., 22: 406-414). Using a different DNA damage detection technology; we tested whether this could be repeated in the nematode Caenorhabditis elegans we also evaluated whether mitochondrial function would be affected. We treated N2 strain (wild-type) nematodes with NNK in liquid culture. Quantitative PCR was applied to analyze NNK-induced nuclear and mitochondria DNA damage. This assay has the advantage of measuring all DNA lesions that inhibit the DNA polymerase, and normalizes results to mitochondrial DNA copy number (Hunter et al, 2010 Methods 51:444-451). Our results confirm that NNK causes both nuclear and mitochondrial DNA damage, but surprisingly nuclear DNA damage was greater than mitochondrial DNA damage in C. elegans. To test whether the mitochondrial DNA damage was associated with mitochondrial dysfunction, we used a transgenic nematode strain that permits in vivo measurement of ATP levels and we used the organic and elemental carbon content (CO and CE, respectively), the presence of microcystine (MN), apoptotic cells (AC) and necrotic cells (NC), and the nuclear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells. The geometric average age was 8.4 years; 41% of children showed at least 1 MN (range 1-6), 27% showed 1-49 AC and 16% had 1-9 clear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells. The geometric average age was 8.4 years; 41% of children showed at least 1 MN (range 1-6), 27% showed 1-49 AC and 16% had 1-9 clear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells. The geometric average age was 8.4 years; 41% of children showed at least 1 MN (range 1-6), 27% showed 1-49 AC and 16% had 1-9 clear division index (NDI) in peripheral lymphocytes by the cytokinesis-block micronucleus test in 1000 cells.

Asbestos is a ubiquitous, naturally occurring mineral fiber that has been linked to the development of malignant and fibrotic lung diseases. Numerous studies have shown that a polymorphism in the genes involved in xenobiotic and oxidative metabolism or in DNA repair processes may play an important role in the pathogenesis of many, if not most, diseases. Unlike some other neoplastic diseases, no association with mesothelioma risk has been consistently demonstrated for polymorphic genes involved with oxidative stress, metabolism, or DNA repair. To evaluate the association between diseases linked to asbestos and genetic variability, we performed a meta-analysis combining the available data on the presence of polymorphisms in GSTM1, GSTT1, NAT2, mEH, XRCC1-C280 and XRCC3-C399T from Italian (n = 57) and Finnish (n = 48) populations with pleural mesothelioma in which the relative magnitude of asbestos exposure was classified. Meta-relative risk statistics were estimated through fixed effects modeling using SAS software. Our analysis found that the GSTM1 null genotype, NAT2 fast acetylator genotype, mEH low activity genotype, and the T/M variant allele of XRC3-C399T were significantly associated with having an increased risk of pleural mesothelioma. When mesothelioma patients were stratified by asbestos exposure (high vs. low), the association between mesothelioma and GSTM1, NAT2, and mEH was not statistically significant for the high exposure group. The association between the T/M variant allele of XRC3-C399T and mesothelioma was significantly increased, regardless of asbestos exposure. The R/Q variant allele of XRC1-R399Q appeared to be confined to those mesothelioma patients with a history of high exposure to asbestos, although the confidence interval includes parity. The results from the present analysis strengthen the hypothesis that intensity of asbestos exposure combined with the presence of certain genetic polymorphisms may increase individual mesothelioma risk.

There are several information on toxicity of textile dyes, but data of colored fabrics are still limited. Although a dye itself may be toxic, its presence in the finished material may not be harmful. Direct Black 38 (DB38) is a benzidine-basedazo dye,

1493 Bariatric Surgery Reduces Intestinal Toxicity in Man.

Obesity is a global epidemic. Worldwide 500 million people are classified as obese with a global prevalence that continues to rise. Obesity and its co-morbidities are amongst the leading causes of global mortality and morbidity and pose substantial socioeconomic burdens on health services worldwide. Bariatric surgery is a form of gastrointestinal surgery that leads to sustained weight loss, resolution of type 2 diabetes and a decrease in cancer risk. These operations alter gut microbial composition which is reflected in the bacterial composition of faeces. We hypothesise that the protective effect of bariatric surgery is due to a decrease in diet-derived intestinal toxic burden. Here, we determined the effect of bariatric surgery on intestinal cytotoxicity and genotoxicity in man. Aquous dimethyl sulfoxide faecal extracts were obtained from preoperative and 2 month postoperative samples from 5 individuals undergoing Roux-en-Y gastric bypass. Human MCL5 cells, a lymphoblastoid cell line expressing CYP1A1, 1A2, 2E1 and 3A4 and epoxide hydrolase, were used to assess faecal extract toxicity and genotoxicity was determined at the TK and HPRT loci using benzo(a)pyrene as a positive control. We found a trend of decreased cytotoxicity postoperatively, as assessed by relative total growth (20% decrease compared to preoperative faecal extracts). Furthermore, the genotoxicity of the faecal extracts decreased postoperatively at both the TK and HPRT loci (50% and 70% respectively, whereas the BaP control induced mutation frequency more than 50 fold). These results support the hypothesis that bariatric surgery leads to a decrease in diet-derived intestinal toxicity burden, which in turn may contribute to the health beneficial effects associated with surgery.

1494 Evaluation of the Mutagenicity of Dyed Fabrics with Direct Black 38.
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There are several information on toxicity of textile dyes, but data of colored fabrics are still limited. Although a dye itself may be toxic, its presence in the finished material may not be harmful. Direct Black 38 (DB38) is a benzidine-based azo dye,
commonly used in textile industry. It is known that, under conditions of perspiration, dyes migrate from colored fabrics and penetrate into human skin. Objective: Evaluate the mutagenicity of DB38 extracted from dyed cotton fibers using artificial sweat solutions. Methodology: Pieces of cotton fabric were dyed with DB38 and washed with and without rinsing (bath with a colloid dispersant). DB38 was extracted from the dyed fabric with artificial sweat solutions (pH 5.5, 6.5, 8.0) at 37°C and 42°C for 2, 8 and 12 h. HPLC-DAD analysis was performed to determine the concentration of dye extracted in each sweat extract. Both original dye and sweat extracts were evaluated by the Salmonella/microsome assay using the strains TA98 and TA100 with and without S9 mix in order to compare with the higher dye concentration extracted with sweat. Results: Original DB38 dye showed positive mutagenic response for TA98 and TA100 with S9. The migration of DB38 dye into artificial sweat resulted in 5.9 μg/mL (dyeing without rinsing, pH 8.0, 42°C, 8 h). Sweat extracts with DB38 did not induce mutagenic effects under the conditions tested. Discussion: Original dye induces mutagenicity by both base-pair substitution and frameshift mutations in presence of S9. However, these effects could no longer be observed after its extraction from dyed fabric using artificial sweat as extracting agent. Our findings showed that DB38 dye can migrate from colored fabrics to artificial sweat, however the concentration of dye extracted does not cause mutagenicity. Thus, as important as the toxicological information of textile dyes, is an investigation of the toxic effects of fabrics which contain these dyes in order to avoid human health problems.

1497 Characterizing the Genotoxicity of the Trichloroethylene Metabolite Dichlorovinyl Cysteine.
J. B. Asfaha, V. De La Rosa and C. Vulpe, University of California Berkeley, Berkeley, CA.

Trichloroethylene (TCE) is an industrial solvent and common environmental contaminant. TCE is metabolized via the glutathione conjugation (GSH) pathway where the reactive metabolite, dichlorovinyl cysteine (DCVC) is formed. Previous work in the field has shown DCVC as the penultimate metabolite resulting in renal toxicity. Our studies using a functional genomics approach in yeast suggest DNA damage and repair pathways play a role in DCVC toxicity. Specifically, we identified the error-prone translesion synthesis (TLS) repair pathway and nucleotide excision repair (NER) as important for response to DCVC exposure. Preliminary work in human fibroblast cells shows initiation of translesion synthesis repair after DCVC exposure. Studies to identify potential DCVC-DNA lesions were conducted using liquid chromatography mass spectrometry (LCMS) analysis. Results indicate the potential for DCVC to cause direct DNA damage via DNA adducts with specific nucleotides. Our data suggests DCVC is genotoxic and has the potential to lead to cell death.

1498 Coexposure to LA Sweet Crude Oil and Dispersant Alters Genomic DNA Damage Occurrence in the Gulf Killifish.
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The BP-Deepwater Horizon event in the Gulf of Mexico unleashed an unprecedented environmental disaster where more than 650 miles of Gulf coastline was littered with Louisiana Sweet Crude oil. During this event Corexit 9500, a surfactant designed to disperse and emulsify crude oil and expedite microbial breakdown, was used to mitigate damage. However, surfactant additives can increase membrane permeability and biological uptake of exogenous compounds including components of crude oil. Recent studies indicate that although visible traces of weathered crude oil have long been absent in the estuarine environments, particular toxic metals persist in organisms and sediments that were once co-exposed to oil and dispersant. This research focuses on the notion that toxic components of crude oil may persist and adversely affect genomic integrity. We examined genomic DNA coding sequences, cytochrome c oxidase-subunit1 and cystic fibrosis transmembrane conductance receptor for lesion analysis. Here we show that killifish, Fundulus grandis, exposed to crude oil and dispersant have increased amounts of genetic damage in both mitochondrial and nuclear templates, relative to killifish exposed to each individual compound. Additionally, DNA lesion formation in the mitochondrial genome was increased in the co-exposures, compared to singular exposures of contaminants.

1499 Absence of Genotoxicity in a Series of 2’-O-Methyl Phosphorothioate Oligonucleotides.

PRO044, PRO045 and PRO053 are 2’-O-methyl phosphorothioate DNA anti-sense oligonucleotides (AON) that are in development for the treatment of Duchenne Muscular Dystrophy (DMD), a lethal orphan disease for which no disease modifying therapy exists. The fundamental cause of this disease is mutations in the dystrophin gene leading to out of frame transcripts for the corresponding muscle protein. AON-induced exon skipping of exon 44, 45 and 53, respectively, in human dystrophin pre-mRNA results in the restoration of the reading frame and an internally deleted but functional dystrophin protein. The mutagenic and clastogenic potential of PRO044, PRO045 and PRO053 was assessed using in vitro tests in bacteria and mammalian cells and an in vivo rodent assay. The bacterial reverse mutation assay (i.e. Ames test using Salmonella typhimurium TA98, TA100, TA102, TA1535 and TA1537) showed that the mentioned AONs have no mutagenic activity at concentrations up to 5,000-10,000 μg per plate, in the presence or absence of an in vitro metabolic activation system (S9-mix). The Chinese hamster ovary (CHO) chromosomal aberration assay showed that PRO044, PRO045 and PRO053 did not induce structural aberrations in the presence or absence of S9-mix, with tested concentration up to 5,000 μg/mL. In addition, PRO044, PRO045 and PRO053 did not induce any polyplody or give indications of mutagenic properties in the presence or absence of S9-mix at 5,000μg/mL. In the in vivo micronucleus assay, no genotoxic effect was observed in the bone marrow of mice following repeated subcutaneous administrations of PRO044, PRO045 and PRO053 at dose levels ranging between 80 and 1,000 μg/kg on days 1, 3, 5, and 8, resulting in a weekly dose of 320 to 4,000 mg/kg. In conclusion, results from these experiments demonstrate that PRO044, PRO045 and PRO053 are not mutagenic nor clastogenic. This is in accordance with the EMA’s reflection paper indicating that phosphorothioate nucleotides (as potential degradation products/metabolites of phosphorothioate oligodeoxynucleotides) are unlikely to pose a genotoxic hazard.

1500 Different Extraction Strategies to Evaluate the Genotoxicity of the Water Soluble Fraction of Air Particulate Matter.
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Exposure to contaminated airborne particulate matter (PM) as a consequence to urban air pollution has adverse effects on human health and the ecosystem. In most air pollution studies, only the genotoxic potential of the organic fraction is evaluated and the genotoxic effects of the soluble metals are not usually verified. In order to determine an appropriate extraction method for assessing the mutagenic potential of the water-soluble fraction of PM, we performed microwave assisted (MW) and ultrasonic bath (UT) extractions, using water as the only solvent, in 10 different air samples (Total particulate matter-TPS- and PM10- fraction of particles with diameter of 10μm or less), from the state of São Paulo, Brazil. Mutagenicity and extraction performances were evaluated using the Salmonella/microsome assay with strains TA98 and TA100, followed by chemical determination of soluble metals (ICP). Except for one sample, the mutagenic potential of TPS was significantly higher in the samples extracted with MW than with UT. The PM10 samples extracted with MW showed mutagenic activity only with TA98, with and without metabolic activation, and in those with UT there was no biological response. Of the metals analyzed (Ni, Cu, Zn, Fe, Pb, Mn, Co, V and Ti), only vanadium (in the samples extracted with MW) and copper (in the UT extracted samples) were above detectable levels. Our results demonstrate the importance in determining an efficient extraction method for assessing genotoxicity, especially at low concentrations of water-soluble contaminants in the air particulate matter. The MW represented a powerful tool to evaluate the genotoxicity of those samples, and additionally has methodological advantages against UT, allowing more reproducibility of the test conditions and improving the analysis precision. Suppprted by CAPES Ph.D. fellowship Isabel Palacio.
mice. A major target for sulphur mustard (SM) is DNA. DNA alkylation leads to the formation of monooadducts, mainly at the N7 position of guanine (HETE-N7Gua: N7 hydroxyethylthioethyl-guanine) and in a lesser extent at the N3 position of ade- nine (HETE-N3Ade: N3 hydroxyethylthioethyl-adenine), and biadducts (N7Gua- ETE N7Gua: bis[N7guanine–ethyl]sulphide). It is a critical step to explain SM cy- totoxicity. Therefore, we have developed a high performance liquid chromatography coupled to tandem mass spectrometry (HPLC-MS/MS) method to quantify the three main SM-DNA adducts. The assay was applied to SKH-1 mice exposed to liquid SM on the dorsal skin at 2, 6 and 60 mg/kg during 4h and were sacrificed at 6h, d1 (day 1), d3, d7, d14 and d21 after exposure. SM exposed skin, skin adjacent to exposed skin and skin being located at 2 cm apart of this latter were removed. The maximum of DNA lesions in SM exposed skin was obtained at 6h post-exposure. The contribution of HETE-N3Ade to SM DNA adducts in skin was much lower than that found in isolated and cellular DNA in previous experiments. Adduct frequencies decreased from d1 likely as the result of cell re- pair. Yet, a relatively important persistence of SM-DNA adducts was observed since HETE-N7Gua and N7Gua-ETE N7Gua were still detected at d21 after 60 mg/kg exposure. For each exposure dose, SM-DNA adducts were also measured, in skin adjacent to the exposed zone, although in lower amounts. They were quantified until d14 after 60 mg/kg exposure. For the highest dose, SM-DNA adducts were detected at 2 cm of the burn but only at d1. These results show a limited radial dif- fusion of SM in skin but a significant persistence in time. They also demonstrate the reliability of our method to quantify SM-DNA adducts which could be moni- tored as part of therapeutic tests.

The OECD Guideline 487 for the In Vitro Mammalian Cell Micronucleus T est (MNvit) became effective in July 2010. The objective of the test is to identify mi- cronucleus formation via clastogenic and aneugenic mechanisms in cells that have completed mitosis. This work presents the assessment of various test parameters for the performance of the assay with human lymphocytes with regard to the above mentioned Guideline. With respect to the In Vitro Mammalian Cell Micronucleus Test (MNvit) aneugenic and clastogenic reference chemicals (colcemid/demecolcin, mitomycin C (MMC) and cyclophosphamide (CPA), were tested for their poten- tial to induce micronuclei in cultured human peripheral blood lymphocytes. The protocol given by the current OECD Guideline 487 was complemented with a modified protocol allowing the cells to have a recovery period between end of treat- ment and addition of the cytokinesis blocker cytochalasin B. Lymphocyte prolifer- ation was studied in response to stimulation with phytohemagglutinin and the nor- mal cell cycle time was assessed by bromodeoxyuridine uptake. The presented data show difficulties in the detection of an aneugenic compounds in binucleate cells and marked differences in the induction of micronuclei in the micronuclei in vitro by the two different test protocols employed by using the four different reference chemicals. Reproducible results were obtained with the clastogenic reference chemical MMC, regardless of the protocol used. With both aneugenic compounds robust and adequate results could be obtained by including the evaluation of mi- cronucleated cells and using the pulse treatment. Robust data were obtained with CPA when using a time schedule including a recovery period.

DNA modifications are considered a trigger for somatic mutations. However, factors such as DNA repair and cell proliferation are known to contribute to DNA modification processes, promoting gene mutations. We previously reported that specific DNA adducts are formed in the livers of rats following treatment with es- tragole (ES), a hepatocarcinogen in rodents. ES caused formation of the DNA adducts, ES-3'-N7Gua, ES-3'-N7dA, ES-3'-N7dG, ES-3'-C8-dG, and ES-3'-N7-DA, in rat livers, as determined by LC-MS/MS analysis. To clarify associations between the number of modified bases and the mutant frequencies (MFs) in the same animal and organ, we quanti- fied ES-specific DNA adducts and performed reporter gene mutation assays in livers, kidneys, and lungs of 6-week-old F344 Sprague-Dawley rats given ES at doses of 0, 3, 30, and 300 mg/kg/day by gavage for 4 weeks. Relative liver and kidney weights were significantly increased in rats given high-dose ES. Mitosis, single-cell necrosis in hepatocytes, and oval cell proliferation were seen in high-dose group livers. Three ES-specific DNA adducts were detected in a dose-dependent manner beginning with the low-dose group. gpt MF was significantly increased only in the high-dose group. In lungs and kidneys, DNA adducts were detected only in the high-dose group, but their frequencies were almost identical to those in the low-dose group livers. These results suggest that the site specificity observed in ES carcinogenicity may depend on the number of modified bases. In addition, the fact that ES-specific DNA adducts were detected to some extent in the livers of rats treated with moder- ate doses of ES, gpt MF being dependent on the involvement of in-vivo-envi- ronmental factors, such as cell proliferation and DNA repair enzymes, in the pro- gression from DNA modification to gene mutation. Some of these factors will be presented, along with results of gpt mutation assays in the lungs and kidneys.

Allura Red AC (Red 40) and Tartrazine (Acid yellow 23) are two of the most commonly used artificial color additives (up to 2% w/w) in food, drug and cosmetic industries. Allura Red and Tartrazine are nitroso derivatives which are the focus of studies on mutagenesis and carcinogenesis because of their transformation into aromatic amines after being metabolized by the gastrointestinal microflora. We in- vestigated the mutagenic activity of Allura Tartrazine in Salmonella typhimurium using the protocol described by Prival and Mitchell (1981). Both TYG104 and TYG1042 strains, derived from TA98 and TA100, respectively carry a plasmid with genes for higher levels of nitroreductase and O-acetyltransferase involved in the break down of azo dyes to mutagenic aromatic amines. These strains have been shown to detect muta- genic activity of azo dyes in the effluents from textile industries, eliciting a higher mutagenic response compared to TA98 and TA100 (Umberto et al, 2003). We used six dose levels (4mM, 2mM, 1mM, 0.5mM, 0.25mM, 0.125mM) of Allura Red and Tartrazine for testing the mutagenic activity. The assay was performed...
under hamster S9-activated and non-activated conditions. In our study, none of the four test strains showed toxicity or an increase in the mutation frequency at the dose levels tested. Whereas, Tartrazine has been reported to have cytotoxic effect in mammalian cells (Patterson and Butler, 1982) and Allura Red has been found to cause DNA damage when tested with the comet assay (Shuij et al, 2001). European Food Safety Authority (2009) has also reported that Allura Red was negative in in-vitro genotoxicity as well as long term carcinogenicity studies. Our results demonstrate that the salmonella strains even with high level of nitroreductase and O-acetyltanferase activities failed to show mutagenic activity in both, Tartrazine and Allura Red.

**1506 Genotoxicity of 2-Bromo-3-Chloropropiophenone.**

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Impurities are generally an integral part of any drug substance or drug product. They can be either process-related impurities that are not completely removed during purification or formed due to the degradation of the drug substance over the product shelf-life. Unlike the drugs, they do not provide health benefits. Therefore, their amount should be minimized. 2-Bromo-3-chloropropiophenone (BCP) is an impurity of bupropion, a second-generation antidepressant and a smoking cessation aid. The US Pharmacoepia recommends an acceptable level for BCP that is not more than 0.1% of the maximum level of bupropion. Recently a qualitative structure-activity analysis (DEREK for Windows version 13) has shown that BCP is likely genotoxic. The genotoxicity of BCP, however, has not been tested based on a search of the available public literature. Because the US FDA has not regulated any genotoxic impurities to much lower levels than non-genotoxic impurities, it is important to determine whether or not BCP is genotoxic. Therefore, in this study the Ames test and the in vitro micronucleus assay were conducted to evaluate the genotoxicity of BCP. BCP was mutagenic with S9 metabolic activation, increasing the mutant frequencies in a concentration-dependent manner, up to 22- and 145-fold induction over the controls in Salmonella strains TA100 and TA1535, respectively. BCP was also positive in the in vitro micronucleus assay, resulting in up to 3.3- and 5.1-fold increase of micronucleus frequency for treatments in the absence and presence of S9, respectively; and 5.9- and 7.4-fold increase of hypodiploids without and with S9, respectively. The addition of N-acetyl-l-cysteine, an antioxidant, reduced but did not eliminate the genotoxicity of BCP in both assays. The results suggest that BCP is mutagenic, clastogenic, and aneugenetic, and these activities are possibly mediated via generation of reactive metabolites.

**1507 Comparison of Mainstream Smoke (Mss) Cytotoxicity and Clastogenicity from Filtered Cigars with the Corresponding Data for the Ms from the Ky34ref Reference Cigarette.**


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Filtered cigars, dimensionally similar to 100s cigarettes, have become popular smoking articles. Unlike cigarettes, the cigarette column is wrapped with a single ply of paper-type reconstituted tobacco. The blends in filtered cigars range from light- to dark-flavored and from mild to strong. Filtered cigarettes are popular with younger smokers, concern has been raised about the toxicity of the smoke relative to cigarette smoke. Rickert et al., 2011, reported that on a per unit smoke nicotine basis the toxicity of the mainstream smoke (MSS) from filtered cigars was no less than that of the MSS from cigarettes. However, the Rickert work used the Health Canada Intensive (HCl) smoking regimen and the MSS from the KY34 reference cigarette. In this research, we have used ISO smoking regimens and three contemporarily 100 mm filtered cigars, one of which (FC1) had a more traditional cigar blend than did the other two (FC2, FC3). On a unit TPM basis, there was little difference in the Neutral Red cytotoxicity of the MSS from the three filtered cigars and the cytotoxicity of the MSS of the KY34ref reference cigarette. However, when the results were put on a per unit smoke nicotine basis, the MSS from two of the three filtered cigars (FC2, FC3), which had low MSS nicotine concentrations, were more cytotoxic than the MSS from the KY34 ref. On a per unit nicotine basis, the cytotoxicity of the MSS from FC1 was similar to that of the MSS of the KY34 ref. Genotoxicity was determined with the in vitro micronucleus assay. On a unit TPM basis, toxicity rankings (−S9) were FC1<FC2<FC3<ky34ref and (−S9) were FC3<FC2<FC1<ky34ref. On a unit nicotine basis, toxicity rankings (both −S9, −S9) were FC3<FC2<FC1<ky34ref. Our results were similar to those reported by Rickert et al. and showed that the ISO smoking regimen was better able to show product differences than was the HCI regimen.

**1508 Effects of Obesity on Spontaneous Reporter Gene Mutations in Gpt Delta Mice Fed a High-Fat Diet.**

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Epidemiologically, obesity has been suggested to be associated with increased risk of several cancers, including cancers of the liver, kidneys, and colon. In several rodent models, genetically induced obesity or high-fat diet (HFD) induced obesity has been shown to enhance carcinogenesis in the liver and colon after treatment with a regimen of respective tumor initiators. However, the factors responsible for the obesity-related progression of carcinogenesis, especially the initiation phase, are not fully elucidated. To reveal the effects of obesity induced by an HFD on spontaneous gene mutations, we performed reporter gene mutation assays in liver, kidney, and colon tissues from obese mice fed an HFD. Six-week-old male and female C57BL/6 gpt delta mice were fed either an HFD or a standard diet (STD) for 13 or 26 weeks. At the end of the experimental period, all mice were sacrificed, and reporter gene mutation assays were performed. Final body weights of male and female mice fed an HFD for 13 weeks were significantly increased compared with those fed an STD, gpt and SpI mutant frequencies in the liver, kidneys, and colon of male mice fed an HFD for 13 weeks were not significantly different from those fed an STD. These results implied that HFD-induced obesity does not influence the spontaneous frequencies of somatic gene mutations, indicating that obesity may affect the tumor promotion phase rather than the tumor initiation phase to enhance carcinogenesis. Further data from the organs of mice fed an HFD for 26 weeks will be presented to evaluate the effects of long-term consumption of an HFD on spontaneous mutagenicity in vivo.

**1509 Community Lead (Pb) Domains and Exposure Disparities: Case Study of Pre- and Post-Katrina New Orleans.**


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We sought to gain insight into the Pb toxicity characteristics of communities in New Orleans. Based on previous soil Pb mapping we divided New Orleans census tracts into two soil Pb domains, high (median ≥100 mg/kg) and low (median <100 mg/kg). Soil samples from four locations represented each of the domains: busy streets, residential streets, house sides, and open spaces (away from streets and houses). Blood and soil Pb concentrations within the high and low Pb domains of New Orleans were analyzed by permutation statistical methods (Multi-Response Permutation Procedures). The children's blood Pb prevalence ≥ 5 μg/dL for the high and low Pb domains were 58.5% and 24.8%, respectively, pre-Katrina vs. 29.6% and 7.5% in post-Katrina New Orleans. In the high Pb domain median soil Pb was 367, 313, 1228, and 103 mg/kg, respectively, busy streets, residential streets, house sides, and open space locations; in the low Pb domain median soil Pb was 64, 46, 32, and 28 mg/kg, respectively (p-Values <0.01-0.16). After Katrina relatively small decreases in soil Pb were observed, and elevated soil Pb permeates the high Pb domain; children living there generally lack Pb safe areas for outdoor play. The low Pb domain was safer by factors ranging from 3 to 38 depending on specific location. Patterns of lead deposition from decades of Pb accumulation have not been transformed by renovations conducted post-Katrina. We expect that all large cities will exhibit the same characteristics as observed in New Orleans. Low Pb soils are available outside of cities to remedy soil Pb contamination. Mapping soil Pb delineates Pb deposition and assists with planning to improve primary prevention of Pb exposure.

**1510 Portable XRF Technology to Quantify Lead and Strontium in Bone In Vivo—Calibration and Validation.**


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Concentrations of lead and strontium in bone can serve as a long-term indicator of exposure levels, and can be more easily correlated to their respective health effects. Lead adversely affects almost all the systems in the body. High levels of strontium, on the other hand, have been correlated to skeletal abnormalities in animal and human populations. We have previously shown the viability of a portable XRF system to quantitate the level of lead in bone in vivo. This system has since been improved to include a larger detector with an improved geometry. In this project, the new instrument was calibrated and validated to quantify lead and strontium in bone in vivo.
The system was effectively calibrated for bone lead quantification with lead doped bone-equivalent phantoms. However, it is difficult to calibrate the system for bone strontium quantification using strontium doped phantoms because most of the bone equivalent materials are severely contaminated with strontium. Therefore, a new calibration method was developed, which made use of a Monte Carlo (MC) simulation model developed specifically for this instrument. In this new method, net strontium signals were obtained for different soft tissue thicknesses using MC simulations and the signals were plotted against corresponding concentrations of the strontium. The concentrations of strontium for in vivo measurements can then be calculated from the calibration lines generated from MC simulation. Three main results were obtained from this study: a) An agreement between the experimental and simulated spectra was achieved; b) the detection limit for bone lead and strontium is improved by a factor of 2 with the improved instrument; c) it is valid to calibrate the system with the calibration lines created by MC simulations. In conclusion, the new system with improved detection limit, combined with the use of Monte Carlo simulations for calibration can be applied to accurately quantify lead and strontium in bone in vivo.

1511 The Effect of Environmental Exposure to Lead on Blood Pressure in Korean Adults.
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General population is exposed to chronic and low level of lead (Pb). Previous studies reported that blood Pb is possibly related to the blood pressure. However, the effect of environmental exposure to Pb on blood pressure is not clear yet. This study was performed to estimate the representative blood Pb level and to assess whether blood pressure is affected by chronic and low Pb exposure in Korean adults. The total study subjects of 2,000 people (male: 908, female: 1,192) were nationwide sampled by multi-staged, sex- and age-stratified probability method, who had not been exposed to Pb occupationally. The geometric mean concentration of Pb in blood was 2.22 ug/dL. The level of blood Pb was significantly higher in males (2.60 ug/dL) than in females (1.97 ug/dL). The level of blood Pb was increased with age-dependent pattern which was the highest in the group of 50-59 years old. Also, blood Pb level was affected by individual behavioral patterns such as smoking and drinking. The mean concentration of blood Pb was higher in rural or urban inhabitants than in metropolitan, respectively. The positive relations were observed between blood Pb concentrations and systolic/diastolic blood pressure in males and in females, both. However, blood Pb was significantly correlated with systolic and diastolic blood pressure after adjustment for covariates in males, but not in females. In summary, the findings from this study suggest that chronic Pb exposure under the environment might be increase blood pressure, systolic and diastolic, both, in males.

1512 Induction of Autophagy and Aberrant MHC Class II Surface Expression in Lead-Exposed Marine Macrophage RAW 264.7 Cells: Dysregulation in Mhc-II Compartment Exocytosis.
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Aberrant major histocompatibility complex class II (MHC-II) surface expression on antigen presenting cells (APCs) is associated with dysregulated immune homeostasis. Lead (Pb) is known to increase MHC-II surface expression on murine peritoneal macrophages ex vivo at concentrations exceeding 25 μM. Little data exist examining this effect at physiologically relevant concentrations. To address this deficit, we examined the effects of Pb on MHC-II surface expression, secondary T-cell activation markers (CD80, CD86, CD40), cell viability, cellular metabolic activity, and β-hexosaminidase activity in RAW 264.7 macrophage cell lines, with changes in cell ultrastructure evaluated by electron and confocal microscopy. Pb induced a bi-phasic, dose dependent increase in MHC-II, CD86, and lysosome-associated LAMP-1 and LAMP-2 surface expression during one cell doubling cycle (17 h), which was mirrored by increased β-hexosaminidase activity. Although cell viability was unaffected, cellular metabolism was inhibited. Electron microscopy revealed evidence of lipid vacuolization, macroautophagy and myelin figure formation in cells cultured with either Pb or LPS. Confocal microscopy with antibodies against LC3B showed a punctate pattern consistent with the presence of mature autophagosomes. Collectively, these data suggest that 2.5 μM Pb increased MHC-II surface expression by inhibiting metabolic activity, inducing autophagy, and increasing MHC-II trafficking in a macrophage cell line.

1513 Exposure to Cobalt Causes Changes in Gene Expression and Protein Abundance in Two Rat Liver-Derived Cell Lines.
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Cobalt is an essential component of the diet, but in large doses it is acutely toxic, and chronic exposures can injure multiple organ systems. While the toxic effects of cobalt have been widely studied, the exact mechanisms of toxicity remain unclear. In order to further elucidate these mechanisms in liver, we exposed two rat liver-derived cell lines, H4-II-E-C3 and MHC1C1, to sublethal concentrations of cobalt chloride to eliminate cell-line specific effects. We chose the treatment concentrations based on a novel qPCR assay using a panel of genes developed from previous metal toxicity studies, as opposed to traditional cytotoxicity assays, to more accurately predict gene expression endpoints at sublethal concentrations. We examined changes in gene expression using DNA microarrays and also examined changes in secreted and cytoplasmic protein abundance using mass spectrometry. We performed both gene-level and functional analyses of the results and observed changes in pathways that are involved in the Nrf2-mediated response, protein degradation, glutathione production, Hif-1 signaling, and energy metabolism. These results are consistent with the known effects of cobalt toxicity, including oxidative stress and hypoxic responses. The changes in protein abundance closely resemble and validate our conclusions drawn from the microarray analysis. This work offers key insights into the role specific genes, proteins, and pathways play in cobalt toxicity mechanisms, and provides leads, which upon validation, may characterize novel toxic effects.

Opinions, interpretations, conclusions, and recommendations are those of the authors and are not necessarily endorsed by the U.S. Army. The research described herein was supported by the U.S. Army Medical Research and Materiel Command, Military Operational Medicine Research Program.

1514 Dose-Response Relationships for Blood Cobalt Concentrations and Associated Health Effects.

Cobalt (Co) is an essential component of vitamin B12. As with all metals, at sufficiently high doses, Co can have detrimental effects on different organ systems, and adverse responses have been observed in patients on Co therapy, workers handling Co-containing powders, and other Co-exposed groups. Although blood Co concentrations are thought to be the most accurate indicator of ongoing Co exposure, little is known regarding the dose-response relationships between blood Co concentrations and adverse health effects. In this analysis, we reviewed the animal toxicology and epidemiology literature to identify blood Co concentrations at which effects have, and have not, been reported. Where necessary, a biokinetic model was used to convert oral doses to blood Co concentrations. Our results indicate that blood Co concentrations in humans of approximately 300 μg/L and higher have been associated with certain hematological and reversible endocrine responses (polycythemia and reduced iodide uptake, respectively), while blood Co concentrations of 700-800 μg Co/L and higher have been consistently associated with a risk of more serious neurological, reproductive, or cardiac effects. Although there are some anecdotal reports to the contrary, the weight of evidence from the many available studies suggest that blood Co concentrations of 300 μg/L and less have not been associated with adverse responses in healthy humans. This suggests that certain populations known to have blood Co concentrations of < 100 μg/L, such as some patients with Co-containing hip implants and those who ingest Co supplements, are unlikely to be at risk due to Co exposure.

1515 Loss of Hypoxia-Inducible Factor (HIF)-1α and Not HIF-2α, in the Lung Alveolar Epithelium of Mice Leads to Enhanced Eosinophilic Inflammation in Cobalt-Induced Lung Injury.
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Cobalt is a known hypoxia mimic and stabilizer of hypoxia-inducible factors (HIFs). HIF-1α contributes to cobalt toxicity in vitro and previous work in our lab has shown that loss of HIF-1α in the alveolar epithelial cells led to a switch from a
neutrophilic to eosinophilic response. While HIF-1α and HIF-2α have known overlapping gene expression targets, HIF-2α is the preponderant HIF isoform in the adult lung, and little is known about HIF-2α in the context of cobalt-induced lung injury. We therefore hypothesized that HIF-2α could be playing a more active role in cobalt toxicity. To compare the roles of HIF-1α and HIF-2α in cobalt-induced lung injury, we used mice deficient in either HIF-1α (HIF-1αΔ/Δ), HIF-2α (HIF-2αΔ/Δ) or both HIF-1α and HIF-2α (HIF-1αΔ/ΔΔHIF-2αΔ/Δ) in alveolar type II epithelial cells. Mice were exposed to cobalt (60 μg/day) or saline via oropharyngeal aspiration for five consecutive days, and sacrificed on day six. Bronchoalveolar lavage (BAL) cellularity, qRT-PCR, and histopathologic analyses were performed. Results confirm previous observations showing that loss of HIF-1α leads to enhanced eosinophilic inflammation, and an enhanced T-helper type 2 (Th2) response. In contrast, inflammation in HIF-2α mice resembled that of control mice following cobalt exposure. HIF-1α/2αΔ/Δ mice showed similar cobalt-induced eosinophilic inflammation seen in HIF-1αΔ/Δ mice. Together, data suggest that loss of HIF-1α in lung epithelium plays the dominant role in determining the response to cobalt-induced lung injury, and that HIF-2α plays a permissive role in this toxicity. Coupled with other experiments performed in our lab, this indicates that epithelial HIF-1α in the postnatal developmental period plays a central role in guiding immune responses in the airway.

1516 Can Zinc Reverse Uranium Toxicity? Potential for a Community-Based Intervention.

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Many Navajo people have been, and continue to be, exposed to uranium through the legacy of uranium mining. 1100 abandoned Cold War uranium sites remain within Navajo communities and numerous wells exceed maximum contaminant levels for uranium and other metals such as arsenic. Certain metals can disrupt protein function by interacting with zinc finger structures and thus inhibit important cellular processes including DNA repair. Based on this mechanism, many metals are now viewed as co-carcinogens and amplify the DNA damaging capacity and tumorigenicty of other carcinogens even at levels where the metals alone are not carcinogenic. The carcinogenicity of uranium is well established in the literature, but there is little known regarding uranium interaction with zinc finger protein structures. Published reports demonstrating that uranium exposure leads to deficiency in DNA repair processes suggest that uranium may interfere with zinc finger DNA repair proteins. Our work with arsenic demonstrates that very low levels of arsenic causes zinc depletion from target zinc finger DNA repair proteins leading to increased DNA damage and mutagenesis that can be reversed by zinc. Based on these findings, we investigated the effect of uranium on DNA repair and the activity of a specific zinc finger DNA repair protein target (PARP-1). Uranium in the form of uranyl acetate (UA) demonstrated little cytotoxicity in an immortalized human embryonic kidney cell line (HEK293) at concentrations at or below 10 μM. UA at concentrations of 10 or 100 μM inhibited the DNA repair protein PARP-1 and caused retention of ultraviolet radiation-induced DNA lesions (CPDs and pH2A.x). The addition of zinc ameliorated PARP-1 inhibition and partially decreased the retention of DNA damage. These findings suggest that one mechanism of uranium toxicity may rely on disruption of zinc finger protein function, so this work will inform a planned assessment of the potential for zinc to block uranium toxicity as an additional component of the Navajo Birth Cohort Study.

1517 Toxicological Properties of PM, from Pellet Combustion—Role of Zn.

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Biomass combustion is important source of ambient air fine particulate matter (PM,10-2.5) and affects human health. Thus, it is important to reveal the effect of used fuel in induced adverse health effects. In search for causative constituents, we studied the role of Zn on toxicity of combustion emission PM10, PM2.5, samples were produced in complete combustion using a pellet boiler (25 kW). Untreated reference pellet fuel (including 11 mg/kg Zn) was used as control sample. The samples were prepared with Zn in different concentrations: 170, 480 and 2300 mg Zn/kg fuel. PM samples were collected on PTFE-filters from diluted flue gas by using Dekati® gravimetric impactor, and subsequently extracted from filters by using methanol. Mouse macrophages (RAW264.7) were exposed for 24 h to four doses (15, 50, 150 and 500 mg/ml) of emission samples. Markers for cytotoxicity (MTT), inflammation (MIP-2 and TNF-α) and cell cycle analysis (DNA content analysis) were measured. All the studied emission samples induced significant dose-dependent decrease in cell viability when compared to control. However, with the samples where Zn was added the toxicity was increased. Moreover, significantly increased in S/G1 phase in the cell cycle was detected after exposure to emissions from all Zn doped pellets, but not with reference pellet. Instead, only minor inflammatory responses were seen by all the samples. The present results indicate that in very good combustion conditions, high Zn content of fuel indicates significant changes in the cell cycle and increases cytotoxicity of PM. Combustion particles toxicity is affected not only by the appliances design or the combustion conditions but also of the fuel and fuel quality.

1518 A Preliminary Study on Analytical Methodology for Determination of Zinc in Liver of Laboratory Rats, by ICP-QMS.

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Zinc is an essential micronutrient for living organisms. It plays a specific role in the synthesis and stabilization of proteins. Its deficiency is characterized by growth retardation, and impaired immune function. While the excess of Zn results in fever and pathological changes in some tissues. Excessive absorption of zinc suppresses copper and iron absorption. Therefore Zn, Cu and Fe can be utilized as pathological indicators. The determination of these elements is of interest in the biomedical area, where laboratory rats are used as model animals, because of their physiological similarities to the human body.

The Inductively Coupled Plasma-Quadropole Mass Spectrometry (ICP-QMS) has been applied to the analysis of trace elements in biological samples due to its characteristics: low detection limit, multielemental analysis and low sample volume.

In this work, analytical methodology was optimized to determine total Zn concentration in livers of laboratory rats. Samples were dried and homogenized, and digested in a microwave oven using HNO3 and H2O2 mixture. Quantitative analysis of the isotope 66Zn using 74Ge as internal standard, was performed using an Elan DRC-e spectrometer (Perkin-Elmer). The dynamic reaction cell was not employed. Results for three replicates of certified reference material (NIST SRM 1577c, National Standard of Standards and Technology) were in good agreement with certified values: 0.21 % (mean relative error, accuracy), and 0.2 % RSD (precision).

The range of Zn total concentrations found in samples of livers of rats were: 37-163 mg/kg. This study belongs to a research where it is included the determination of Zn/Cu ratio in several tissues. The authors thank the financial support of DGAPA-PAPIIT IN229911 and PAL 3400-02.

1519 Effects of Tungsten Chemical Species on Biochemical Pathways and Mineralization in Bone.

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Tungsten (W) is a metal that has numerous civil and military applications. When W enters the environment, it is rapidly oxidized and speciates based on the environmental matrix it is embedded in. Bone is the long-term storage organ for W—and presumably W chemical species if present—when taken up by organisms. It is unknown what long-term effects W has in bone, therefore, we evaluated the effects of W species on biochemical pathways in hFOB 1.19 osteoblastic cells exposed to water (control), or W chemical species (sodium tungstate, phosphotungstate [PW], poly tungstate [polyW], and tungstosilicic acid [TSA]) at 0-10,000 μM. Preliminary analyses also demonstrate an effect on alkaline phosphatase, the enzyme involved in bone mineralization. Synchrotron analysis of bone from female rats exposed to 200
mg/kg/d sodium tungstate in drinking water ad libitum for 90 d showed W incorporation as calcium poly-tungstate (77% total W in bone) and calcium tungstate (23%). These data demonstrate that W chemical species generally only affect phosphate-dependent cell signaling and secondary messenger pathways in hFOB 1.19 cells, and may also affect bone mineralization enzymes, resulting W species incorporation into bone in animals. Follow-up studies will analyze the effects on bone mineralization and strength associated with W incorporation due to chronic exposure.

1520 Accumulation of Manganese (Mn) in Rat Brain Ventricular Region following In Vivo Subchronic Mn Exposure: Effect on Copper (Cu) Status.

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Welders and smelters occupationally exposed to high levels of Mn have an increased risk for developing manganism, a disease similar to Parkinson's. Existing human and animal studies indicate that Mn accumulates mainly in the striatum, the area adjacent to the brain ventricular region (BVR) which houses cells implicated in adult neurogenesis. The neural stem cells in BVR are capable of migrating to striatum and hippocampus en route to the olfactory bulb. This group has previously shown Mn exposure results in Mn accumulation in the choroid plexus (CP), cerebrospinal fluid (CSF) and increases intracellular Cu concentration. This study was undertaken to test the hypothesis that a high Mn level in the CP and CSF may influence the surrounding ventricular structure and interfere with the homeostasis of other essential elements. Adult male rats received daily ip injections of MnCl2 at the doses of 6 and 15 mg Mn/kg as the low and high dose groups, respectively, or saline as the control, 5 days per week for 4 weeks. Twenty-four hr after the last injection, BVR, femur bone and liver tissues were dissected and quantified by atomic absorption spectroscopy for concentrations of Mn, Cu, iron (Fe) and zinc (Zn). Data analysis by Student's t-test showed that levels of Mn in BVR of the low- and high-dose groups increased by 94.8% and 218.5%, respectively, as compared to controls (n=4-6, p <0.01). In femur, Mn concentration increased by 278.2% and 1237.4% in the low- and high-dose groups, respectively (p < 0.01). Interestingly, Cu concentrations in BVR decreased significantly for both low and high-dose groups (p<0.05). In contrast, Cu levels in bone increased by 23.2% and 26.5% in the low- and high-dose groups, respectively ( p<0.05). Taken together, our data provide the initial evidence that Mn accumulates in brain ventricular area following in vivo exposure. Mn's potential impairment of this region in the disease progression of Mn Parkinsonism deserves further investigation. (Supported in part by NIH RO1 ES088146)

1521 Methylmercury-Induced Dopaminergic Neurotoxicity in Caenorhabditis elegans.

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Mercury is a persistent environmental contaminant that exerts its toxic effects on the nervous system through molecular mechanisms that remain unknown. Parkinson's disease (PD) is a neurodegenerative disorder characterized by the slow progression and irreversible loss of the dopaminergic (DAergic) neurons. The etiology of PD remains elusive, with aging, environmental toxicant exposure and genetic mutations contributing to the disease. Epidemiological studies have pointed to the contribution of methylmercury (MeHg) to DAergic vulnerability and the predisposition to PD. We examined the impact of early-life exposure to MeHg on DAergic neurodegeneration in Caenorhabditis elegans (C. elegans) with emphasis on gene-environment interactions. SKN-1, orthologue of mammalian Nrf2, is a major stress-activated cytoprotective transcription factor. We hypothesized that MeHg's toxicity is dependent on an intact SKN-1 response and skn-1 knockout (KO) worms would show heightened toxicity compared to wildtype (N2). We also tested the effect of MeHg in a pdr-1KO worm, ortholog to the human parkin gene (mutated form found in juvenile PD), under the premise that these worms would show heightened sensitivity to MeHg. We identified the impact of early-life MeHg exposure on MeHg content, stress reactivity and measures of DAergic neurodegeneration in N2, skn-1KO and pdr-1KO worms exposed to 0-30μM MeHgCl2 for 30 minutes following synchronization. Our data suggests that skn-1KO (LD50=19μM) and pdr-1KO (LD50=17μM) are more sensitive to MeHg than N2 controls (LD50=25μM). MeHg uptake was higher in the pdr-1KO strain compared to the N2 strain. DAergic morphology observed via fluorescent analysis at adult life-stage revealed presence of puncta at 20μM MeHg in skn-1KOs and loss of cell body fluorescence in pdr-1KO. Dopamine (DA)-dependent behavioral analysis revealed alterations in DA following MeHg exposure, corroborated by decreasing DA levels, motor tone and levodopa uptake as well as decreased tyrosine hydroxylase (TH) activity. Taken together this data suggests that exposure to MeHg early in the lifespan can have detrimental effects on DAergic neurons. Supported by R01ES07371 and EST2007028.

1522 Lactational Exposure to Mercury (Hg) and Evidence of Oxidative Stress.

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The objective of this work was to assess exposure to Hg in mothers and their breast-fed infants. We also evaluated urinary 8-hydroxy-2-deoxyguanosine (8-OHdG) and malonaldehyde (MDA) in both the mothers and infants as biomarkers of oxidative stress. The mean concentration of Hg in breast milk was 1.19μg/L (range:0.012-6.44μg/L) with only one mother had Hg<4μg/L, the ATSDR upper limit. This corresponds to approximately a daily intake of 1.02μg Hg (0.13-5.47μg/kg). Based on the average infant's body weight in this study (7.7kg) and the EPA reference dose of Hg (0.3 μg/kg/day), this brings the highest daily intake to 2.31μg/kg. There were 7 infants that exceeded this limit. None of the mothers had total blood Hg > the EPA reference dose of 5.8 μg/L. Probably, this might be due to the low exposure to organic in inorganic Hg. Furthermore, no correlation was noted between UHg in infants and Hg in breast milk (P=0.05). On the other hand, Hg in breast milk was associated with total Hg in blood (r=0.53, P=0.001) suggesting the efficient transfer of organic form of Hg from blood to milk. Both urinary MDA (r=0.2, P=0.015) and 8-OHdG (r=0.24, P=0.003) levels increased with the mother's UHg levels. Infant's UHg levels in infants were positively associated with urinary levels of MDA (r=0.31, P=0.003) and 8-OHdG (r=0.43, P=0.00) in infants. There was also an association between Hg in breast milk and increased urinary MDA levels in infants (r=0.35, P=0.0). Our results clearly showed the transfer of Hg from the mother to infant through breastfeeding, and high exposure induces oxidative stress. Breastfeeding can be a potential health risk. Nevertheless, breastfeeding should not be discouraged and instead efforts should be made to identify and eliminate the source of Hg exposure in the present population.

1523 Methylmercury Toxicity in KK-Ay Obese Type 2 Diabetic Mice.

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We examined the toxic effects of methylmercury (MeHg) exposure on the brain, kidney, pancreas and spleen of KK-Ay type 2 diabetic mice to clarify how the metabolic changes associated with type 2 diabetes mellitus (T2DM) affect MeHg toxicity. MeHg (5 mg Hg/kg/day p.o.) was given to male KK-Ay and C57BL/6 (BL/6) mice three times per week for 6 weeks beginning at 4 weeks of age. Average of body weights (BW) of vehicle-treated BL/6 and KK-Ay mice were 16.3 g and 16.4 g on the first day, and 24.8 g and 42.3 g on the last day of the experiment, respectively. BW gain in MeHg-treated KK-Ay mice stopped 4 weeks after MeHg administration, and these mice began to lose weight around 5 weeks. Six of 7 MeHg-treated KK-Ay mice showed neurological symptoms such as hind limb crossing in the final stage of the experiment. Alternatively, no obvious symptoms and decrease in body weight were observed in vehicle-treated and MeHg-exposed BL/6 mice. The mean blood mercury level of KK-Ay mice administered MeHg increased constantly throughout the administration of MeHg and reached a maximum of about 8 μg/mL. On the other hand, the average blood mercury level of BL/6 mice administered MeHg was 2.8 μg/mL after 10 days of MeHg administration. The average kidney weights of vehicle-treated and MeHg-exposed KK-Ay mice were 542 mg and 408 mg, and those of BL/6 mice were 357 mg and 413 mg, respectively. The average spleen weights of vehicle-treated and MeHg-exposed KK-Ay mice were 91 mg and 135 mg, and those of BL/6 mice were 65 mg and 71 mg, respectively. The average total mercury concentrations in the cerebrum, kidney, pancreas, spleen and epididymal fat pad of KK-Ay mice were 27, 51.7, 40.9, 32 and 1.6 μg/g, respectively. The average total mercury concentrations in these organs of BL/6 mice were 7.4, 10.4, 6.9 and 0.57 μg/g, respectively. These results indicate that body fat gain in T2DM and low accumulation of mercury in adipose tissue can increase organ accumulation of MeHg accompanied by toxicity in KK-Ay mice.
1524 The Palmitoylation of Meh1, a Component of EGO Complex, Has an Important Role in the Reduction of Methylmercury Toxicity in Budding Yeast.

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Methylmercury (MeHg) is an environmental pollutant that causes the severe central nervous system disorder. However, the molecular mechanism underlying MeHg toxicity remains to be clarified. We have found that deletion of the Akr1, one of palmitoyl transferase, causes hypersensitivity to MeHg in budding yeast. In this study, we found that yeast cells lacking Meh1, one of substrates for Akr1, exhibited hypersensitivity to MeHg. Palmitoylation of Meh1 was not detectable by a deletion of Akr1, suggesting that Akr1 is the main palmitoyl transferase for Meh1. We also found that the palmitoylation of Meh1 by Akr1 is essential for the protective effect of Meh1 against MeHg toxicity. Meh1 has been identified as a component of the EGO complex that is involved in the regulation of mitoautophagy. We found an increased susceptibility to MeHg in yeast cells with deletion of Meh1 or the other EGO complex components (Gtr2 and Ego3). Moreover, Meh1/Gtr2- or Meh1/Ego3-double deletion mutants did not exhibit further increase in the susceptibility to MeHg when compared to Meh1 deletion mutant. These results suggest that Meh1 might reduce MeHg toxicity as a component of the EGO complex. It has been reported that Meh1 forms an EGO complex by binding with Ego3 on vacuole membrane. Interestingly, Meh1 with a mutation in the palmitoylation site was defective in its ability to bind with Ego3. Moreover, normal Meh1-GFP was localized in the vacuolar membrane, but the mutant Meh1-GFP was widely distributed in the cytoplasm and not in the vacuolar membrane. These phenomena proposed that the palmitoylation of Meh1 has an important role in its cellular distribution and formation of the EGO complex.

As for our result, the palmitoylation of Meh1 by Akr1 is related to the formation of the EGO complex through its localization into the vacuolar membrane, and this distribution and formation of the EGO complex is important for the reduction of MeHg toxicity.

1525 Involvement of AAT Transporters in Methylmercury Toxicity in Caenorhabditis elegans.

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Caenorhabditis elegans (C. elegans) is a powerful genetic model for the investigation of neurotoxicity of metals, such as methylmercury (MeHg). MeHg has been shown to accumulate in C. elegans, leading to decreased survival, reproductive and developmental defects, and neurodegeneration. It is unknown how MeHg is transported in the worm. In mammals, MeHg complexes with cysteine, which functionally resembles methionine; and the complex is transported into cells via a molecular mimicry mechanism involving the large neutral amino acid transporter LAT1. It is unknown if MeHg can enter cells without being complexed to cysteine or if there are additional uncharacterized MeHg transport mechanisms. C. elegans can be used to quickly identify MeHg transport mechanisms due to their ease of genetic manipulability and high degree of homology to mammals. Herein we tested the hypothesis that LAT1 homologues (AATs) are responsible for MeHg transport in C. elegans. Worms were pre-treated with 1 mM methionine to assess whether MeHg toxicity could be attenuated by increasing a competitor for AAT transporters. Treatment with MeHg resulted in decreased survival of C. elegans (LD50 = 22.91 μM), however methionine pre-treatment showed significantly less toxicity to MeHg (LD50 = 47.54 μM, p < 0.0001), implicating the AAT amino acid transport systems in MeHg toxicity. Using RNAi feeding protocols we selectively knocked down expression of the aat-1, aat-2 and aat-3 genes, and assayed for MeHg induced lethality. Knockdown of each gene individually resulted in a significant right hand shift in the dose-response lethality curve (aat-1: LD50 = 118.1 μM, p < 0.001; aat-2: LD50 = 102.1 μM, p < 0.001; aat-3: LD50 = 44.65 μM p < 0.001), suggesting these genes may be involved in MeHg transport. As lethality of MeHg was completely blocked by methionine pre-treatment or aat-3 gene knockdown, there is a possibility that additional MeHg transport mechanisms may be identified in C. elegans.

1526 Effects of Methylmercury on Heart Rate Variability in the Rat.

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Analysis of heart rate variability (HRV) in humans has revealed that methylmercury (MeHg) exposure decreases parasympathetic nerve function or increases sympathetic nerve function. However, limited information is available on such effects in experimental animals. In the present study, 5 rats were treated 2 times per week for 5 weeks with MeHg (4 mg/kg) or vehicle orally. Several weeks before and after treatment, telemetry recordings of ECG were acquired. HRV and its power spectrum parameters were determined from 1-hour segments (approximately 16,000 to 20,000 beats) of ECG data during the day by LabChart and its HRV Module. With 4 mg/kg MeHg treatment, a decrease in body weights and the hind limb circumference and an increase in the heart rate (HR) were observed from the analysis of the ECG. HRV analyses showed a decrease in the coefficient of HR (CVVR) and low frequency (LF) and high frequency (HF) HRV. A slight increase in the LF/HF ratio was observed due to the decrease in HF. These results suggest that a decrease in the CVVR, LF, and HF and an increase in the HR indicate the suppression of parasympathetic nervous system activities after repeat treatments with MeHg in rats. These findings also suggest similar effects of MeHg on the autonomic nervous system of humans and animals.

1527 Sex Differences in Paraaxonase Activity in Subchronic Inorganic Mercury Exposure.

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Epidemiological evidence suggests an increase of cardiovascular diseases (CVD) as a result of inorganic mercury exposure. The underlying mechanisms underpinning this risk, as well as sex differences in response to mercury exposure, are not yet understood. Paraaxonase (PONase), an enzyme located in the high-density-lipoprotein (HDL) has been shown to protect against CAD. In order to investigate the association between inorganic mercury exposure and CAD, male and female rats were exposed to 0.5, 1.0 and 1.5 mg/kg HDL-PON activities towards paraoxon (PONase) and phyanylacetate (AREase) in plasma, lipoproteins, hepatic and brain microsomal fractions were determined using standard methods. Inhibition of PONase and activation of AREase in plasma and HDL characterised the effects of inorganic mercury in both sexes. Inorganic mercury exposure inhibited PONase by 63% (plasma) and 67% (HDL) respectively in male animals, whereas the female enzyme was inhibited by 80 and 47% respectively. AREase activity was activated by 55 and 53% in male, whereas the activation in female amounted to 25 and 49% respectively. In the VLDL, inorganic mercury inhibited PONase in both sexes whereas AREase was activated in female animals but inhibited in male. In the hepatic microsomal fractions, only the PONase enzyme was inhibited in male animals whereas in female, activation was observed in both enzymes at the highest dose of inorganic mercury. Brain microsomal cholesterol was increased in male but decreased in female by inorganic mercury resulting in altered cholesterol/phospholipid ratios. Our findings indicate that inorganic mercury exposure exerts an inhibitory effect on PONase but activated AREase. Modulation of PON activity may be an early biochemical step in the induction of CAD by mercury. This may also be mediated through changes in membrane fluidity brought about by changes in the concentration of cholesterol in the microsomes.

1528 Ethylmercury Induces ER Stress and Mitochondria Dysfunction Mediated Autophagy.

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Methylene mercury is one of the most important environmental and industrial pollutants throughout the world. Exposure to mercury causes strong damage to organs including the brain, blood, liver, bone and kidneys. Renal proximal tubular cells represent the primary target site. We previously investigated the cytotoxicity of seven kinds of mercury compounds in human renal proximal tubule (HK-2) cells. Ethylmercury chloride (EMC) was shown the strongest cytotoxicity among them with 2.4 and 0.76 μM of IC50 values exposed for 24 h and 48 h in HK-2 cells, respectively. In this study, we explored the mechanism of EMC induced cytotoxicity in HK-2 cells. EMC was increased the accumulation of mtDNA from 0.5 to 2 μM concentrations in a dose-dependent manner. The expression of MT-I and Hic-5, which were related to oxidative stress, was also induced by EMC treatment. In addition, calcium is a well-known regulator of many intracellular processes, including apoptosis and autophagy. Intracellular [Ca2+] was increased in cytosolic [Ca2+], which was the tendency appears unlikely that the increase of MT-I and Hic-5, which were related to oxidative stress, was also induced by EMC treatment. In addition, calcium is a well-known regulator of many intracellular processes, including apoptosis and autophagy. Intracellular [Ca2+] was increased in cytosolic [Ca2+], which was the tendency appears unlikely that the
male C57BL/6 mice with 1, 2, 5 and 10 mg/kg/day. The expression of ER stress related marker genes were increased by immmunohistochemical analysis. ICSH-II had significantly increased in the kidneys of EMC treated mice. Together, these results suggest that EMC induced autophagy via ER stress and mitochondria dysfunction not only in human renal proximal tubule (HK-2) cells but also in mice kidneys.

1529 Protective Role of Quercetin against Mercury-Induced Nephrotoxicity in Sprague-Dawley Rats.
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Mercury chloride (HgCl2) is a well-known nephrotoxicant. HgCl2 preferentially accumulates in the kidneys and causes nephrotoxicity as a result of oxidative stress. The aim of this study was to investigate the protective role of the antioxidant activity of HgCl2-induced acute renal damage. Quercetin, a naturally occurring phenolic compound found in fruits and vegetables, has gained considerable attention for its antioxidant properties. In this study, quercetin (250 mg/kg) was administered orally to Sprague-Dawley male rats for 3 days prior to HgCl2 (20 mg/kg) treatment. All animals were sacrificed 24 h after the last treatment and various urine or blood biomarkers associated with renal damage were measured. Significant increases in urinary volume and decreases in urinary pH were observed in the HgCl2-treated group. Blood urea nitrogen (BUN) and serum creatinine (sCr) levels were significantly increased in the HgCl2-treated group when compared to control group. In the HgCl2-treated group, the urinary excretion of BUN, sCr, lactate dehydrogenase (LDH) and total protein were all markedly lower than those of the control group. These adverse effects were ameliorately by pretreatment with quercetin. Compared to the HgCl2-treated group, quercetin pretreatment led to a significant decrease in the amounts of KIM-1, NAGL, HMGB1, and Netrin-1 in the urine. Furthermore, histopathological examinations indicated that quercetin had a protective effect against HgCl2-induced proximal tubular damage. These results suggest that quercetin may be a promising agent to protect against HgCl2-induced acute kidney injury.

1530 Selenium Protects against the Toxic Effects of Methylmercury in Sperm Whale (Physeter macrocephalus) Skin Cells.
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Mercury (Hg) persists, bioaccumulates and is toxic to the marine environment posing risks to the marine ecosystem. Selenium (Se), an essential element, antagonizes toxicity of Hg by increasing glutathione levels and the activity of glutathione peroxidase (GPx) in the kidneys of rats exposed to Hg. The aim of this study was to determine if Se protected sperm whale skin cells from Hg-induced toxicity at similar concentrations found in the skin biopsies. Sperm whale skin cells developed from the skin biopsies were cotreated with 72 h exposure to 2, 3, 4 or 5 μM methylmercury (MeHg) and 10 μM sodium selenite, and cytotoxicity, mitotic index and genotoxicity was measured. Our findings show that MeHg induces cytotoxicity, alters mitotic index and causes aneuploidy in sperm whale skin cells, and Se protects against MeHg-induced toxicity compared to control.

1531 Selenium Compounds Cause Apoptosis and GSH Depletion in Rat Hippocampal Astrocytes.
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Selenium (Se) is a required trace element in mammalian systems. While generally accepted that active work play in a protective role, exposure to high levels of Se have been associated with neurotoxicity. Previous work in our laboratory has demonstrated toxicity of Te and Se compounds in rat hippocampal astrocytes. The purpose of this study was to evaluate the mechanism of cell death of diphenyl diselenide (DPDSe) and selenium tetrachloride (SeCl4) and to investigate their effects in the functioning of mitochondrial complexes III and IV and depletion of the antioxidant GSH. Rat hippocampal astrocytes were used as a model system. The LC50 of both compounds had been previously found to be 7.8 μM. For the present study, concentrations of both compounds ranging from 9.15-625 μM were tested. Phase-contrast microscopy was used to study the morphological changes in the cells following treatment. Micrographs of cells exposed to high concentrations of either compound (15.625 and 7.8μM) showed decreases in cell number and increases in rounded cells with loss of processes. Micrographs of cells exposed to low concentrations (3.9 and 1.9μM) appeared similar to control astrocytes. Inductively Coupled Plasma-Optical Emission Spectrometry (ICP-OES) demonstrated that treatments with both compounds resulted in a dose-related increase of Se in the cells that is significantly greater than amounts seen in the controls. Caspase 3/7 activities were assessed and significant increases in activity were observed at the 7.8 μM and 15.625 μM concentrations with both Se compounds confirming apoptotic activity. ELISA assays were used to investigate the activities of complexes III and IV of the mitochondrial respiratory chain. There were no significant changes in the activities of the two enzymes in any treatment group. GSH/GSSG ratios were significantly decreased at 7.8 and 15.625 μM of each compound indicating oxidation of sulfhydryl groups. These results suggest that selenium compounds cause cytotoxicity in rat hippocampal astrocytes resulting in apoptosis but mitochondrial complexes III and IV are not involved.

1532 Heavy Metal Hazards of Smokeless Tobacco in Nigeria.
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Smokeless tobacco (ST) or snuff prepared by grinding tobacco leaves into a very fine powder with addition of salt petre and other ingredients has been in use in Nigeria for a very long time. ST is marketed for oral and nasal use. Recently there has been an upsurge in the use of ST even amongst the younger generation. Information on the public health implication and heavy metal hazards of ST in Nigeria is sparse. This study was carried out to determine the extent of heavy metal contamination in ST used in Nigeria. Concentration of five heavy metals, Lead(Pb),Cobalt(Co),Cadmium(Cd), Nickel(Ni), and Chromium(Cr) were determined in smokeless tobacco samples from six geographical regions in Nigeria. Our results showed that the level of contamination with all the metals was ≤10μg/g, 11-30μg/g,≤10μg/g, ≤20-70μg/g and ≤270-11400μg/g for Pb, Co, Cd, Ni and Cr. These concentrations are higher than acceptable limits. Health risks associated with ST snuffing have received little or no attention in Nigeria in spite of wide spread use. This study underscores the importance of elaborate human risk assessment of ST in Nigeria and indeed Sub Saharan Africa.

1533 Tobacco Cigarettes: A Source of Metals (Al, Cd, Co, Cr, Mn, Ni, Pb, Sr) for Humans.
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Every six seconds a smoker dies, the victim of an addiction that causes a high degree of toxicity. Al, Cd, Co, Cr, Mn, Ni, Pb and Sr were determined in 33 tobacco samples randomly obtained in Tenerife using Inductively Coupled Plasma Spectrometry by previously treating the samples according to the standard procedure, that is, by adding HNO3 at 65% to the sample and burning it in a muffle oven for 50 hours at 450 degrees Celsius. With respect to results, 428 mg Al/kg, 0.810 mg Cd/kg, 0.558 mg Co/kg, 1.442 mg Cr/kg, 112.026 mg Mn/kg, 2.238 mg Ni/kg, 0.602 mg Pb/kg and 82.206 mg Sr/kg. Al was the metal with the greatest concentration while Co had the lowest. The tobacco was also classified according to the type of tobacco (Virginia vs Dark), type of cigarrette (Normal vs Light) and the manufacturers confirming apotic activity. The amount of Al in Atlantic Virginia tobacco was 423 mg/kg while in Dark this was 478 mg/kg, Cd concentrations were 0.807 mg/kg for Virginia and 0.844 mg/kg for Dark, Co values were 0.56 mg/kg for Virginia and 0.533 mg/kg for Dark, Cr values were 1.487 mg/kg for Virginia and 1 mg/kg for Dark, Mn was determined to be 113.362 mg/kg for Virginia and 98.667 mg/kg for Dark, Ni values were 2.242 mg/kg for Virginia and 2.199 mg/kg of each compound indicating oxidation of sulfhydryl groups.
for Dark, Pb afforded 0.607 mg/kg for Virginia and 0.556 mg/kg for Dark, and Sr gave values of 77.982 mg/kg for Virginia and 124.444 mg/kg for Dark. Regarding the differences between Light and Normal tobacco, the following values were obtained in Light: 412.4 mg Al/kg, 0.873 mg Cd/kg, 0.553 mg Co/kg, 1.64 mg Cr/kg, 120.02 mg Mn/kg, 2.4 mg Ni/kg, 0.873 mg Pb/kg and 78.627 mg Sr/kg; while in Normal tobacco the values determined were 434.783 mg Al/kg, 0.783 mg Cd/kg, 0.559 mg Co/kg, 1.366 mg Cr/kg, 108.56 mg Mn/kg, 2.168 mg Ni/kg, 0.484 mg Pb/kg and 83.762 mg Sr/kg. Exposure to metals though the inhalation of cigarette smoke represents a significant source of metal contamination to smokers, adding to the dietary intake of these same metals.

**1534 Metal Mobilization from Retained Embedded Fragments in a US Veteran: Biomonitoring Correlates with Fragment Content.**

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Recent clinical findings implicate the potential for local and systemic long-term health effects from embedded metal fragments associated with injuries from improvised explosive devices in soldiers serving in military conflicts. In the past, fragments embedded in muscle tissue were thought to be relatively inert; however, evidence has shown that soldiers with embedded depleted uranium fragments have elevated urine U levels 20 years after exposure. To understand the potential health risks associated with embedded fragments from blast injuries, the Department of Veterans Affairs has established a medical surveillance program which integrates fragment composition data, surrounding tissue analyses, and urine biomonitoring results that characterize systemic and local tissue exposure to: Al, As, Cd, Cr, Co, Cu, Fe, Mn, Ni, Pb, U, W and Zn. These metals were chosen based on available fragment composition data and known toxicity of individual metals. We present here results from a Veteran who had 3 fragments removed several years after injury. Using EDXRF, the removed fragments were determined to be an Al-Cu alloy. Chemical analysis of adherent tissue showed levels of Al and Cu consistent with mobilization of these elements from the fragment to the surrounding tissue. Histology showed focal foreign body-type giant cell reaction and chronic inflammatory cell infiltrates, but no evidence of neoplastic changes. Prior to fragment removal, biomonitoring results showed a urine Al level 1.5 fold higher than the reference range, but below levels associated with adverse health effects. Concentrations of all other metals were within established reference ranges. These results provide further evidence that metal fragments in the body are not inert and materials released from fragments can enter systemic circulation over time, thus warranting long-term biomonitoring and medical surveillance of Veterans with embedded fragment contents. Supported by the Department of Veterans Affairs.

**1535 Baseline Blood Levels of Manganese (Mn), Lead (Pb), Cadmium (Cd), Copper (Cu), and Zinc (Zn) in Residents of Beijing Suburb.**

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The baseline blood concentrations of heavy metals are important for monitoring metal exposure in the general population. The data can be used to evaluate the current environmental pollution and also serve as the references for exposure assessment in local residents as well as in occupational settings. The purpose of this study was to determine the blood levels of Mn, Pb, Cd, Cu, and Zn among the residents (aged 12-60 years old) living in the suburb southwest of Beijing, China. Blood samples were drawn from a total of 648 subjects from March 2009 to February 2010. Metal concentrations in the whole blood were measured by ICP-MS. The geometric mean of the blood levels of Mn, Pb, Cd, Cu, and Zn was 11.4, 42.6, 0.6, 802.4 and 4.665 µg/L, respectively. Blood Mn level of all participants is similar to the reports by Korean (Lee et al. Int J Hyg Environ Health 2012;215:449) and Canadian (Clark et al. Chemosphere 2007;70:155) studies in literature. Blood Pb level was higher in Chinese residents than those of Korea and Czech (Batarova et al. Int J Hyg Environ Health 2006;209:359). Blood Pb levels in the all participants were higher among Chinese than those of Korean and Czech; the magnitude of increase became more evident when compared with Korean data. Blood Cd levels in this Chinese cohort were within the same range as Czech, but were significantly higher than German’s (Heitland & Koster J Tissue Elem Med Biol 2006;20:253). Blood Cu levels were significantly higher in Chinese than those of Germany and Canadian, while blood Zn levels were significantly lower than those of Czechs. Based on the data of the 95th percentile, we estimate the normal reference values for the whole blood Mn to be 21.7 ng/mL, Pb 10.3 µg/dL, Cd 5.3 ng/mL, Cu 1.36 µg/mL, and Zn 9.03 µg/mL in Chinese resident living in Beijing suburb. These baseline values will be useful for occupational and environmental exposure assessment.

**1536 Effect of Long-Term Exposure of Heavy Metals on the Expression Profile of Cytoprotective Genes among Individuals Living in Mining Areas.**

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Environmental exposure to heavy metals is one of the most global concerns, particularly among whom living around the polluted areas. These heavy metals are known to generate oxidative stress species, and induce of oxidized DNA adduct formation, and DNA repair genes causing disease induction. The aim of this study is to evaluate the effects of long-term exposure to environmental heavy metals in the expression profile of some cytoprotective genes and the level of oxidized DNA adduct formation among individuals living around mining areas. Sixty healthy volunteers were divided to two groups: exposed group consisted of 40 male residents in a heavy metal-polluted area, whereas the control group consisted of 20 male residents in a non-polluted area. Whole blood concentrations for heavy metals, particularly lead, cadmium, and mercury, were determined using ICP-MS. Total RNA was isolated using PAXgene Blood RNA kits and the mRNA levels of heavy metals biomarker genes; NQO1, HO-1, and MT-1 and DNA repair genes: OGG1 and APE1 were determined by RT-PCR. Plasma levels of 8-OHdG were determined by ELISA technique. The results showed a significant increase in blood levels of heavy metals among exposed group. RT-PCR results revealed significant inhibitions of NQO1 mRNA level in group which were highly contaminated with lead, whereas the group of highly contaminated with cadmium showed the most inhibitory effect on MT-1 mRNA level and significant induction of OGG1 mRNA level. Furthermore, mercury contaminated group showed a significant inhibition of APE1 mRNA level. The level of 8-OHdG was well correlated with the level of OGG1 mRNA. We conclude that long-term exposure to environmental heavy metals differentially altered the expression of cytoprotective and DNA repair genes. Therefore, they may be used as biomarkers for early prediction of disease risk among individuals living around polluted areas.

**1537 Phytotoxicity and Trace Metal Accumulation from Long-Term Chicken Litter Amendment in Wilkes County, North Carolina: A Field Study.**


Arsenic, zinc and copper are commonly integrated in chicken feed to ensure increased weight gain in chickens and prevent chicken diseases. These metals ultimately become part of the litter that is spread by farmers over crop fields as an alternative to expensive fertilizers. Massive growth in industrial chicken production across Wilkes County, NC has resulted in increased concentrations of phytotoxic metals in farmland soils and loss of consumer sensitive crops such as peanuts. Using Zea mays as a representative crop species, our research objective was to evaluate the long-term sustainability of this practice by measuring adverse effects on corn fields that have received a varied number of poultry litter amendments. Three agricultural fields receiving chicken litter amendment ranging from less than 10 yrs to more than 30 yrs within Wilkes County were selected as sampling areas for trace metal analysis. Soils were collected prior to and after annual litter application dates and corn tissue samples were collected in thirty day intervals (30, 60 & 90 days post planting) from each field. Nearly 400 corn tissue and soil samples were lyophilized and homogenized; 0.5g samples were nitric acid digested with a microwave reactor (USEPA Method 3051); and analyzed using ICP. OES to quantify levels of Cu and Zn. ICP- OES analysis indicated that industrial broiler chicken litter contains considerably elevated Cu (~388 ppm) and Zn (~481 ppm) concentrations. Thirty day corn seedling Cu and Zn burdens correlate closely with soil metal levels and with significantly delayed seedling emergence time and reduced growth rates (e.g., mean leaf height and area). No correlation was observed between soil metal levels and metal concentration in chicken litter. Continuous litter application may result in significantly reduced Zea mays productivity due to elevated Cu and Zn levels and therefore is not sustainable at the maximum application rates currently practiced in NC.
In the wake of the Deepwater Horizon oil disaster of 2010, we collected tissue samples from sperm whales in order to determine the effects of the oil and chemical dispersants on their population. We focused on whales because they are at the top of the food chain and they are biologically similar to humans. Sperm whales live and breed in the waters surrounding Torreón, Mexico, where we collected our samples. The levels of metals in the whale samples were compared to the levels found in crude oil and dispersants. The results showed that the levels of metals in the whale samples were similar to those found in the oil and dispersants. This suggests that the dispersants are not significantly altering the metal levels in the whale tissues. The findings have important implications for the health of the sperm whale population and for the environmental impact of oil spills in the future.
to 5.4 or 54.1 mg/m³, respectively) did not result in any adverse findings and show a clear threshold of response for lung and liver. Therefore, the no-observed-adverse effect concentration (NOAEC) was determined to be 10 ppm (equivalent to 54.1 mg/m³).

### 1543 Evaluation of Nose-Only Inhalation Exposure to Aerosolized Isoeugenol in Sprague-Dawley Rats.

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Isoeugenol is a commonly used fragrance material that has been associated with skin sensitization. The study objective was to understand the effects of inhalation exposure in Sprague-Dawley (SD) rats when aerosolized when administered to rats by nose-only inhalation at 0.15, 1.5, and 14.9 ppm (1, 10, and 100 mg/m³) for 2 weeks (6 hours/day, 5 days/week). Prior to the main study, a preliminary 3-day exposure (6 hours/day) was run at 89.4 and 44.7 ppm to allow selection of appropriate doses. At both concentrations, the animals were observed to have decreased body weight over the 3 day exposure period. Therefore, a third of the lowest dose tested, 14.9 ppm, was chosen as the high exposure level for the main study. Standard endpoints evaluated included: clinical observation; body and organ weights; hematologic and serum chemistry evaluation; macroscopic/microscopic examination of selected organs; and bronchoalveolar lavage fluid (BALF) and serum analysis for cellular markers of inflammation (i.e., cytokines). As a positive control to validate the utility of cytokine measures as indicators of pulmonary inflammation, a nose-only inhalation pilot was conducted with 25 mg/m³ amorphous silica to elicit a chronic inflammatory response. In both exposure groups, there were no toxicologically significant effects on BALF or serum cytokine levels. Therefore, neither a no-observed-adverse-effect level (NOAEC), nor a no observed effect level (NOEL) could be determined for this study regarding upper airway irritation. For the lower airway the NOEL was considered to be 14.9 ppm.

### 1544 Toxicological Evaluation of 4, 4, 5, 5, 6, 6, 7, 7, 8, 8, 8-Undecfluoro Octanoic Acid (5:3 Acid).

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Studies were conducted to evaluate mammalian and aquatic toxicity of 4,4,5,5,6,6,7,7,8,8,8-undecfluoro octanoic acid (5:3 acid) (CAS# 914676-49-3). 5:3 acid is a metabolite of 6:2 fluorotelomer alcohol (6:2 FTOH). 6:2 FTOH is a raw material for fluorotelomer-based products. Rat oral and dermal LD50 values were 2950 mg/kg and >5000 mg/kg, respectively. 5:3 acid produced no dermal irritation but irreversible eye irritation in rabbits, while a weak dermal sensitization response was observed in mice (LLNA), with an EC3 of 23%. A bacterial reverse mutagenicity assay and an in vivo rat micronucleus assay were negative. A 2-week rat gavage study identified red blood cells (anemia), liver (hypoproteinemia, focal necrosis), and increased beta-oxidation, kidney (tubular vacuolation), stomach (ulceration), and tongue, kidneys, thymus, and testes were collected, weighed, and evaluated histologically; serum testosterone (T) and hepatic CYP 2A5 (equivalent to human CYP2A6 in humans and the major nicotine and cotinine oxidase in mouse liver) was also measured. Heart weight (relative to body weight) was significantly decreased following either nicotine or gutkha exposure, while normalized liver/body weight and serum T levels were significantly decreased in the Gutkha-treated group only. These findings suggest that repeated Guthkha use adversely impacts body weight, organ weight, and circulating T levels and that Guthkha toxicity may be driven by toxic components other than nicotine. As the use of Guthka rapidly increases worldwide, future studies should further elucidate its toxicological implications.

### 1545 A 28-Day Toxicity Study of PSOA by Oral Gavage in Rats Followed by a 14-Day Recovery Period.

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The objective of this study was to determine the potential repeat dose toxicity and reversibility of any findings over a 14 day recovery period for a persoy sulfurated fatty acid (PSOA). The study design followed OECD Guideline 407. Sprague-Dawley (CrlCD(SD)) rats were given PSOA once daily via gavage for 28 dates at doses of 0, 5, 15 and 50 mg/kg/day. The highest PSOA concentration used was 0.5%. On completion of the dosing period, designated recovery animals were retained for a 14 day recovery period. The following parameters and end points were evaluated in this study: clinical signs, body weights, body weight changes, food consumption, ophthalmology, functional observations, clinical pathology parameters, gross necropsy findings, organ weights, and histopathology. No early deaths were observed in the dose groups. At the end of the treatment period, gross and microscopic findings observed were of the nature and incidence commonly seen in rats of this age and strain for studies of this type, and/or were of similar incidence in control and treated animals. Differences in the incidence of dark foci in the lungs, observed in male rats sacrificed on Day 29, were shown to correlate with alveolar hemorrhage which showed no biologically significant difference in incidence between treated animals and controls. The increased incidence of minimal hemorrhage in the thymus of treated animals which were sacrificed after the recovery period on Day 43 is considered incidental, as this is a common background lesion. Overall, no test article-related findings were observed for this study. Also no impairment of the mucous membranes of the gastrointestinal tract was observed up to the highest concentration of 0.5%, which can be regarded as a No-Observe-Effect-Concentration (NOEC) in terms of local toxicity after 28 days repeated exposure. In conclusion, administration of PSOA by once daily oral gavage was well tolerated in rats at levels of 5, 15 and 50 mg/kg/day. Based on these results, the No-Observed-Effect Level (NOEL) was considered to be 50 mg/kg/day.

### 1546 Smokeless Tobacco (Gutkha) Induced Toxicological Effects in a Mouse Model: A Nicotine Twist.

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The popularity of smokeless tobacco (ST), usually placed within the mouth to be chewed, sucked, or swallowed, is growing rapidly and its prevalence of use is rising globally, due (in part) to greater convenience and stricter smoking laws. Gutkha, an addictive form of ST that contains areca nut, catechu, cardamom, lime, flavored chewing tobacco, and natural and artificial flavoring materials, is particularly common amongst Southeast Asian communities throughout the world, including the US. This study seeks to determine the generalized toxic effects of Guthkha and to determine the role of nicotine in producing Gutkha-associated toxicity. Ten-week-old B6C3F1 male mice were divided randomly into three groups and treated for 3 wk (5 d/wk) via the oral mucosa with equal volumes of water (control), a water-soluble nicotine solution, containing 0.24 mg of nicotine, or a water-soluble, 21 mg hypophosphat Guthkha solution. Serum cotinine, used as the exposure metric, was measured weekly in all three groups and was similar in both the Guthkha- and nicotine-treated mice (36 vs. 48 ng/mL, respectively). At sacrifice, liver, heart, spleen, tongue, kidneys, thymus, and tests were collected, weighed, and evaluated histologically; serum testosterone (T) and hepatic CYP 2A5 (equivalent to human CYP2A6 in humans and the major nicotine and cotinine oxidize in mouse liver) was also measured. Heart weight (relative to body weight) was significantly decreased following either nicotine or gutkha exposure, while normalized liver/body weight and serum T levels were significantly decreased in the Guthkha-treated group only. These findings suggest that repeated Guthkha use adversely impacts body weight, organ weight, and circulating T levels and that Guthkha toxicity may be driven by toxic components other than nicotine. As the use of Guthka rapidly increases worldwide, future studies should further elucidate its toxicological implications. MSK Cancer Center and NIEHS Center Grant ES000260.

### 1547 Safety Assessment of Dicamba Monooxygenase from the Biotechnology-Derived Soybean MON 87708.

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Monsanto has developed a soybean, MON 87708, which is tolerant to dicamba herbicide. It contains a gene, dmo, that expresses dicamba monooxygenase (MON 87708 DMO). A safety evaluation of the MON 87708 DMO protein was conducted, it examined: 1) the source of the gene; 2) its function and a history of safe use for structurally similar proteins; 3) its homology to known allergens, toxins, or...
other proteins known to have adverse effects in mammals; 4) its stability to heat.

**1548** A 14-Day IV Infusion Exploratory Toxicity Study of Hydroxylamine in Rats.


Hydroxylamine (HA) is an important industrial reducing agent with little therapeutic value. However, it may be formed as a by-product when drugs containing hydroxamic acid functional groups are used, and thus exhibit hematotoxicities. As many of these compounds are developed as intravenously (IV) administered (infusion) drugs, understanding the dose-toxicity relationship of HA by IV infusion becomes important in risk mitigation and management. In this study, we investigated if, at the same daily doses (12, 24 or 36 mg/kg/day), either by continuous infusion (CI) or by twice daily 1-hour intermittent infusions (II), HA produced the same level of toxicity. As systemic toxicities of HA are well-known, the focus of this study was on toleration and hematologic effects. Clinical signs, body weight and quantitative food consumption were assessed. Blood samples were taken at various times to evaluate methemoglobin (metHb) and total hemoglobin (Hb) levels. Hematology parameters and morphology were evaluated in blood samples taken on Days 3, 9, 15 and 30. Dose-dependent increases in the metHb level were observed in both CI- and II-treated animals, however, a higher level of metHb was generally observed in II animals at the same daily doses. The metHb level increased within the first 4-6 days of dosing and then declined in the remaining course of study. Clinically, administration of HA up to 36 mg/kg/day (CI or II) was well-tolerated, with only palpebral cyanosis observed in the high-dose animals (CI and II) and not in the historical controls. The second week. No significant effect of HA on the body weight was observed although food consumption at 36 mg/kg (CI and II) was significantly reduced. Multiple hematology parameters demonstrated HA induced a dose- and time-dependent hemolytic anemia with the formation of Heinz bodies in both CI and II animals. The reduction of total Hb worsened with daily dosing and appeared to depend upon hemolytic anemia with the formation of Heinz bodies in both CI and II animals. The reduction of total Hb worsened with daily dosing and appeared to depend on the formation of Heinz bodies in both CI and II animals. The reduction of total Hb worsened with daily dosing and appeared to depend on the formation of Heinz bodies in both CI and II animals.

**1549** Increased Hypothalamic Dopaminergic Neuron Tyrosine Hydroxylase Expression in Lean Wistar Rats.

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Caloric restriction in Wistar rats has been reported to show a decrease in mammary and pituitary tumours and an increase in uterine tumours (1, 2) associated with delayed hypothalamic dopaminergic neuronal senescence (3). This study was designed to assess the effects of reduced body weight gain on hypothalamic dopaminergic neurons using FPEP tissue obtained from 100 female Wistar rats (50 controls; 50 with >30% body weight gain reduction) from a 2 year bioassay. A combination of immunostaining (IHC) and RNAAscopeTM in situ hybridisation (ISH) was used to examine expression of tyrosine hydroxylase (TH), a rate limiting enzyme for dopamine synthesis. The TH protein expression was detected in both the paraventricular (PVH)/periventricular (PeVH) nuclei and the median eminence (ME). TH RNA expression was also detected in these regions, but there was a disparity in the strength and location of TH RNA expression compared to the protein expression. TH protein staining was strong in the PVH/PeVH nuclei and was not associated with signs of toxicity. Taken together, dietary exposure to intact/active MON 87708 DMO is anticipated to be negligible. MON 87708 DMO was also administered to mice in acute and 28-day toxicity studies, and the protein was not associated with signs of toxicity. The weight of evidence demonstrates that MON 87708 DMO is safe for food and feed.

**1550** Proposition 65 and Cancer Incidence in California.


Proposition 65 has been in effect in California (CA) for 25 years. It requires businesses to notify consumers if products they buy contain chemicals that are known to the State of CA to cause cancer and birth defects or other reproductive harm. This study determined the tumor types reported to be associated with the 71 carcinogens that were added or removed to the Proposition 65 list in 1987 and evaluated the incidence rates for these tumor types in males (M) and females (F) in CA through to 2009. Corresponding incidence rates in Washington State (WA); geographically similar) and New York State (NY) as a diverse East coast regional control) were evaluated as comparators. Of the 71 chemicals added as carcinogens to the Proposition 65 list in 1987, >10 have each been reported to be associated with urinary bladder and breast tumors and >20 each with liver and lung tumors. Between 1988 and 2009, there was a modestly decreased trend in bladder cancer incidence in CA [linear slope = 0.19 (M) and 0.06 (F); R2 = 0.75 (M) and 0.77 (F)]. Less robust decreases were observed in WA [slope = 0.05 (M and F); R2 = 0.01 (M) and 0.18 (F)], while a markedly increased trend was observed in NY. Over the same time period, there was a modestly increased trend in breast cancer incidence [slope = 0.15 to 0.41; R2 = 0.02 to 0.18] and an approximate doubling of liver cancer incidence in both sexes in all 3 states [slope = 0.07 to 0.41; R2 = 0.80 to 0.99]. A substantial and comparable decrease in lung cancer incidence was observed in males in all 3 states [slope = 1.18 to 1.80; R2 = 0.90 to 0.99]. For females, lung cancer incidence decreased in CA (slope = 0.56; R2 = 0.79) but increased in WA and NY. The results of this preliminary assessment indicate that since Proposition 65 came into effect 25 years ago, liver cancer incidence in CA has approximately doubled but consistent and robust decreases in incidence rates have been observed for bladder cancer in both sexes and for lung cancer in females. The contribution of Proposition 65 to these observed changes is unclear and warrants further investigation.

**1551** Alternatives to Bisphenol A in Thermal Paper: Analysis of Substitution Options and Trade-Offs.


The U.S. Environmental Protection Agency (US EPA) Design for Environment (DfE) Program undertook a chemical alternatives assessment for the use of bisphenol A (BPA) in thermal paper, including cash register receipts, as part of the Action Plan for BPA in March 2010. The purpose of the alternatives assessment is to identify and compare alternative chemicals to inform decision-making. The alternatives assessment was conducted via a multi-stakeholder partnership that identified 19 functional alternatives to BPA. The hazard assessment for BPA and the alternatives used the DfE hazard evaluation criteria to assign hazard designations for human health toxicity, ecological toxicity and environmental fate endpoints. Some alternatives were well characterized for all endpoints. Other alternatives were poorly characterized, wherein analog data, predictive models, structural alerts and expert judgment were used to make hazard designations for data gaps. Trends for human health, ecological toxicity and fate characteristics were indicated in a number of the alternatives due to particular molecular size ranges and/or molecular structures. In general, all alternatives have important trade-offs although if used with appropriate control measures, some of the alternatives could provide incremental benefits. Environmental hazard assessments, coupled with decision-making protocols that are practical tools for businesses to use in materials selections, will lead to more sustainable product development when human health or ecological toxicity concerns exist. The resulting hazard profiles should be of value to manufacturers making substitution decisions and facilitate reductions in environmental releases and subsequent exposures.
Personal care product (PCP) ingredients are released into aquatic ecosystems via wastewater discharge and potentially affect environmental health. The environmental safety assessment (ESA) is an important component of overall ingredient safety evaluation. We presented here an exploratory effort in applying the body-residue based approach in ESA of PCP ingredients. The traditional water-concentration based ESA has been used to evaluate the environmental impact of PCP ingredients. Compared with the water-based concentration, body residue concentration represents an internal dose, takes chemical disposition into account and provides an integrated assessment of the exposure an organism receives over space and time. With advancement of computational toxicology, critical body residues (CBRs) corresponding to toxicity endpoints and chemical bioaccumulation levels can be modeled as an alternative to animal testing. Organic UV filters are widely used in PCPs to offer sunscreen benefit. Environmental occurrence of various organic UV filters has been documented in both abiotic and biotic media. As a lipophilic UV filter, benzophenone-3 bioaccumulates in fish and concerns have been raised due to its in vitro activity. The goal of this study was to explore the possibility of a threshold level for ocular irritation from repeated topical exposure to agents in solution.

**Methods:** Ten compounds from 8 chemical families (acids, acrylates, alcohols, aldehydes, amines, anionic surfactants, and cationic surfactants) were identified which resulted in severe ocular irritation or corrosive effects when administered as a single drop on the rabbit eye at high concentrations. Each chemical was prepared in an appropriate vehicle (saline or sesame seed oil) to give concentrations of 20 and 100 ppm. One eye of albino rabbits (5/dose group) was administered 50 μL of the test article 4 or 6 times daily (oil- or saline-vehicle, respectively) for 3 days. The contralateral eye served as a vehicle control. Draize scoring was performed before the first daily dose and after the final daily dose and biomicroscopy was performed pre-dose and on Days 2 and 3.

**Results:** None of the 10 chemicals resulted in notable ocular irritation at either 20 or 100 ppm after repeated ocular administration.

**Conclusion:** Despite being severe irritants or corrosives when dosed as a single drop at high concentrations, the favorable ocular irritation results obtained with up to 100 ppm concentrations of severe irritants or corrosives suggest that it may be possible to establish a threshold for ocular irritation.

**1554 Human Health Assessment of Scented Candle Emissions.**

Airborne compounds in the indoor environment arise from a wide variety of sources such as environmental tobacco smoke, heating & cooking, dusts, emissions from furniture and construction materials as well as outdoor sources. One product category which has received recent attention as a source of indoor airborne substances is scented candles. The potential impact of airborne candle emissions on the quality of the indoor air is highly dependent on the type and concentrations of chemical substances released. To better understand the potential of scented candles to contribute to the indoor load of airborne substances, a comprehensive candle emission testing program initiated by the consumer products and fragrance industry was undertaken to investigate the emissions of volatile and semi-volatile organic compounds (VOC; SVOC) and particulate matter (PM). Associated human exposures scenarios were derived and computer models used to estimate exposure of these materials to consumers. Measured chamber concentrations of VOC, SVOC and PM were used to predict their respective, cumulative indoor air concentrations in a standard EU-based dwelling using 2 models - the well-known and widely accepted ConsExpo 1-Box inhalation model and the recently developed, refined RFID 2-Box indoor air dispersion model. The output from both models has been used to estimate realistic yet conservative consumer exposure to scented candle emissions measured under this program. The potential consumer health risks associated with the exposure to these materials was compared to existing air quality guideline values and established safe exposure levels. This investigation concluded that even under the conservative assumptions, potential human exposures are a minimum of one order of magnitude below established regulatory indoor air guideline values and/or published safe exposure levels.

**1555 Cobalt Whole Blood Concentrations in Healthy Adult Volunteers Following Two Weeks of Ingesting a Cobalt Supplement.**

Recently, there has been an increase in the marketing and sales of dietary supplements, energy drinks, muscle builders and other consumer products that may contain relatively high concentrations of essential elements. Cobalt-containing supplements are readily available in the U.S. and have been marketed to consumers as energy enhancers. However, little information is available regarding cobalt (Co) body burden and steady-state blood concentrations following the intake of Co dietary supplements. We assessed Co whole blood concentrations in four healthy adult male volunteers who ingested a commercially available Co supplement (0.4 mg Co/day) for 15 or 16 days. Pre-supplementation blood Co concentrations were less than the reporting limit of 0.5 μg/L. Consistent with background Co concentrations reported to range between 0.2 to 0.4 μg/L. The mean whole blood Co concentration in the volunteers after 15 or 16 days of dosing was 3.6 μg Co/L and ranged from 1.8 to 5.1 μg Co/L. The mean observed concentration in the study group was approximately 9 to 18 times greater than background concentrations. Further studies of Co whole blood concentrations following supplementation over longer time periods with additional monitoring of physiological parameters may provide useful information for evaluating the health of persons who take various doses of Co.

**1556 Multiparameter In Vitro Toxicity Testing of Crizotinib, Sunitinib, Erlotinib, and Nilotinib in Human Cardiomyocytes.**

Targeted therapy has greatly improved the treatment and prognosis of multiple types of cancer. However, unexpected cardiotoxicity has arisen in a subset of patients for some of the tyrosine kinase inhibitors (TKi). For these TKi, the cardiotoxicity was not wholly predicted by pre-clinical testing, which centers around the inhibition of the human Eater-a-go-go-Related Gene (hERG) channel. Therefore, we sought to determine whether a multi-parameter panel of tests that assesses a drug’s effect on cellular, molecular, and electrophysiological endpoints would more accurately predict cardiotoxicity. To do so, we examined how 4 FDA-approved drugs impacted cell viability, apoptosis, reactive oxygen species (ROS) generation, metabolic status, impedance, and ion channel function in human cardiomyocytes. The 3 drugs with known associated cardiac adverse events (crizotinib, sunitinib, and nilotinib) all proved to be cardiotoxic in our series of in vitro tests while erlotinib, a cardiac-safe drug, did not show any indications of toxicity. Crizotinib, an ALK/MET inhibitor, was the most cardiotoxic by our panel, leading to increased ROS production, caspase activation, cholesterol accumulation, and a significant disruption in cardiac cell beat rate and blockage of ion channels. The multi-targeted TKi sunitinib also demonstrated severe cardiotoxicity in our tests, showing decreased cardiomyocyte viability, inhibition of AMPK, increased lipid droplet accumulation, disrupted beat pattern, and hERG block. Nilotinib, a second
generation Ber-Abl inhibitor, led to increased ROS generation, caspase activation, hERG block, and an arrhythmic beat pattern. Thus, each drug showed a unique toxicity profile, demonstrating that a multi-parameter approach allows for a more complete assessment of the potential for drug-induced cardiotoxicity and may allow for earlier detection in the drug development process.

1557 Safety Assessment of Chemicals in Toys.
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Numerous countries have enacted laws and regulations to protect children from the potential health hazards associated with exposures to chemicals used in the manufacture of toys, including the United States, Canada and the member states of the European Union. Assessing the risk posed to children from exposure to chemicals in toys poses a special challenge, due to the particular vulnerability of this population to both adverse effects and chemical exposure. Examples of chemical categories used in toys that may be of concern include metals, boron-containing substances, pigments and colorants, preservatives, and allergenic fragrances. The aim of this study was to evaluate the risk of substances flagged as potential chemical hazards when included as ingredients in child-intended products in various jurisdictions. Results of toxicological studies indicate that metal-containing pigments, certain azo dyes, and boric acid and its salts are associated with toxicity concerns such as reproductive, developmental, mutagenic, and carcinogenic effects. Product formulation data collected in a proprietary database over a two-year period was analyzed in order to determine the levels and frequency of use of these substances in various categories of toys. Exposure considerations and scenarios designed to specifically address per and for children from chemical substances as a result of infants’ inten-
tional use and reasonably anticipated misuse of different categories of toys were developed. Using the results of the toxicological studies and exposure scenarios developed by our group, we analyzed the margin of safety for the use of these chemicals of concern with respect to toxicity effects in toys. The results of our studies suggest that restricted chemicals are still being used to formulate toys at levels that indicate potential concern for this sensitive population.

1558 Health-Based Framework for Evaluating the Safety of Hydraulic Fracturing Products.
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Hydraulic fracturing has made it possible to extract natural gas from dense shale rock formations and has become the fastest-growing source of gas in the U.S. Because the process involves drilling through groundwater formations, there has been concern that drinking water aquifers could become contaminated—a concern compounded by a lack of information regarding the composition and safety of the products used. In an effort to address these concerns, we developed a quantitative framework to characterize potential safety of hydraulic fracturing products. The framework consists of four evaluation criteria that are applied to each of the product components: composition and use, toxicity, exposure, and risk of release. Each of the criteria has specific requirements associated with scores ranging from 1-5 (with 1 being the best and 5 being the worst) that are based on USEPA guidelines and standard risk assessment practices. A final composite product score is calculated based on scores for each of the criteria for each of the components and this score is used to place the product into a category of use relating to its safety (not recommended for use, use with caution/specialized conditions, or acceptable for designated use). Importantly, if any component of the hydraulic fracturing product does not achieve a score below the STOP point for any of the four criteria (indicative of a minimum level of knowledge or safety), the product is automatically placed into the not recommended for use category. This framework allows health experts to tailor the evaluation to any type of product used in the fracturing process and can be consistently conducted by independent parties. Because this process focuses on human health risk and environmental exposures, it is different than those currently available in the natural gas industry and is more in line with standard risk assessment practices. Most importantly, this framework allows for transparency in the evaluation process and provides a quantitative method upon which experts can make decisions.

1559 Novel Methodology in Hazard Assessment: Chemical Clustering for Read Across—The Phthalate Alternatives Case Study.

Phthalates are a class of compounds produced in high volume in the U.S. and are found in many products, primarily as plasticizers. The U.S. EPA published an initial Action Plan for eight phthalates in 2010 due to concern about the toxicity of phthalates and the evidence of pervasive human and environmental exposure. As indicated in the Action Plan, a Design for the Environment (DfE) alternatives assessment (AA) will be performed for these chemicals. The DfE Program publishes AAs to help industries identify safer chemicals and provide a comparison of potential human health and environmental impacts of chemical alternatives. DfE hazard criteria are used to assign hazard designations for human health toxicity, ecological toxicity and environmental fate endpoints. Over 70 substances were identified as potential alternatives to the eight action plan phthalates. Some alternatives are well characterized for all endpoints, whereas others are data poor. In the absence of experimental data, DfE assessment methodology designates hazards based on a read across approach to structurally similar compounds. This poster describes a novel technique for green chemistry and hazard screening that uses the EPA’s Office of Pollution Prevention and Toxics ChemACE program to cluster the alternatives based on common structures, functional groups, and molecular architectures. The ChemACE program automates chemical clustering based on structural similarities and generates reliable and organized results. Using this methodology, read across (analog) data from data-rich chemicals was used to assist in assigning and justifying hazard designations for data poor chemicals within a cluster. This allows for the determination of hazard designations for as many endpoints as possible including human health endpoints that often lack experimental data and methods for estimation. The resulting AAs should be of value to manufacturers making substitution decisions and facilitate reductions in potential human health impacts.

1560 Transdermal Toxicity of the Phorbol Ester Isolated from Biodiesel Feedstock, Jatropha Curcas.
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[Purpose] Jatropha curcas attracts rising attention as a biodiesel feedstock in the world. However, Jatropha contains various toxic components, generating concerns about its health effects. One of toxic components is potential tumor promoter, phorbol esters. Toxicity of Jatropha phorbol esters has not been fully described. In this study, transdermal toxicity of main component of Jatropha phorbol esters was assessed in mice.

[Experimental procedures] One of the Jatropha phorbol esters, 12-deoxy-16-hydroxyphorbol-4-[(12’,14’-butadienyl)-6-[(16’,18’,20’-nonatrienyl) bicyclo[3.1.0]hexane-(13’-O)-2’-[carboxylate]-(16’)-O]-3’-[8’-butoenoic]-1’-10’ (DHPB) (1 – 10 μg), was applied onto skin of mice. As a positive control, 1,2,3,4- Tetradeanoylphorbol 13-acetate (TPA) (1 – 10 μg) a known tumor promoter was applied. Transformation activity of test compound was assayed using Bhsa42 cells. Autopsy was conducted on the dead mouse to observe the lesions. All survived mouse were sacrificed at 8 weeks from the beginning of the study and hematological and splenic lymphocyte measurement using flow cytometer were carried out.

[Results] 2.5 μg of DHPB led to weight loss and over 5 μg of DHPB caused death during 5 weeks observation. There were no such symptoms in the mice treated with the same dose of TPA. Gastrointestinal bleeding and splenic atrophy was observed in the dead mice. The NOAEL of DHPB was considered to be 2 to 2.5 μg. No papilloma on the skin was observed in DHPB group in contrast to TPA which developed papilloma when the dose exceeded 1 μg. There was no significant difference in splenic lymphocyte composition among DHPB, TPA, and control groups. Hematological analyses showed that the number of red blood cells, platelet, and lymphocyte was markedly decreased in DHPB treated mice at 8 weeks.

[Conclusion] DHPB, on the contrary to TPA, showed no tumor promotion in this experiment but showed acute toxicity which was not seen in the mice treated with TPA. These results suggest that DHPB has different characteristics in transdermal toxicity in comparison with those of TPA.
A key quantitative element in toxicity studies is the identification of the NO(A)EL – no observed (adverse) effect level. With the introduction of omics into toxicology, questions have been raised concerning the sensitivity of these methods. BASF’s Experimental Toxicology and Ecology unit and metanomics developed the MetaMap® tox database with more than 500 reference compounds obtained from rat studies (OECD 407 design). Metabolome analysis in plasma was performed after 7, 14 and 28 days and relative levels of endogenous metabolites in treated rats versus controls were analyzed. We have obtained metabolome data at toxicological NO(A)EL doses.

Recent advances in metabolic profiling technologies together with expert judgment offer the possibility of identifying whether differences from control values are treatment-related and to discriminate between those that are adverse and those that are not. To obtain a measure of sensitivity of metabolomics vs. classical toxicology, we have used these data to analyze metabolomics changes at toxicological NO(A)EL doses. We have done this considering the number of statistically significant metabolite changes (p<0.05), the false-positive rate and the correlation with defined patterns for diverse modes of action (approximately 100 defined in MetaMap® tox).

Results show that in most cases where there are no toxicological effects (NOEL) there are also only few metabolomics changes (at/below the level of the false positive rate and without a match to the predefined MoA patterns). In some cases in which the study demonstrated a NOAEL (only effect noted being liver weight increase) metabolomics changes are often present and identify the liver as target organ. Following this analysis, it would seem that metabolomics is generally not more sensitive than classical toxicology, with respect to the identification of a NOAEL.

Metabolic Profile of Rats in Repeated Dose Toxicological Studies after Oral and Inhalative Exposure.

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BASF and metanomics, in a joint effort, developed the MetaMap® tox-application containing metabolic profiles attained from administration of over 500 toxicants. The toxicants were administered to five Wistar rats for each sex and each of the two dose levels for a total duration of four weeks following a standardized protocol, in accordance with OECD 407. Metabolome analysis of plasma samples was performed after one, two and four weeks. The ratios of the metabolites from treated rats versus untreated rats (control) were transferred to MetaMap® tox for data assessment.

In this study, metabolic profiles after oral and inhalative exposure were compared for six selected model compounds: aniline (A), chloroform (CL), ethylbenzene (EB), 2-methoxyethanol (ME), N,N-dimethylformamide (DMF) and tetrahydrofuran (THF). The toxicants were dosed inhalatively for six hours/day, five days a week and orally each day via the feed (DMF) and by gavage (A, CL, EB, ME, THF). Statistical evaluation (pairwise comparison in MetaMap® tox as well as principle component analyses) showed similar metabolic changes for DMF, EB, CL and ME, indicating a comparable systemic toxicity after oral and inhalative exposure. In contrast, metabolic profiles of A and THF showed differences between the tested exposure routes. Metabolic profiles were sufficient for the identification of test substance related toxicological modes of action using MetaMap® tox. In conclusion, our results indicate the potential of metabolic profiling for generating fingerprints of systemic toxicity following different exposure scenarios.
The Barnett Shale is a large natural gas reserve encompassing more than 5,000 square miles and covering approximately 26 counties in North Texas. Due to public concern that natural gas compressor stations emit high formaldehyde concentrations, formaldehyde has been of increased public and regulatory interest in recent years. It has also been detected in 24-h carbonyl samples collected by the TCEQ in

Recently, medical research has seen a strong push toward translational research, or “bench to bedside” collaborations, which strive to enhance the utility of laboratory science for improving medical treatment. The success of that paradigm supports the potential application of the process to other fields, such as risk assessment. Close collaboration among academic, government, and industry scientists may facilitate the application of scientific findings to regulatory decision making. The National Toxicology Program (NTP), National Institute of Environmental Health Sciences (NIEHS), and U.S. Food and Drug Administration (FDA) developed a consortium-based research program to more effectively link academic and guideline-compliant research. An initial proof-of-concept collaboration, the Consortium Linking Academic and Regulatory Insights on BPA Toxicity (CLARITY-BPA), uses bisphenol A (BPA) as a test chemical. The CLARITY-BPA program combines a core perinatal guideline-compliant 2-year chronic toxicity study with mechanistic

The Endocrine Disruptor Screening Program (EDSP) has been designed to determine whether certain substances may have effects on the estrogen, androgen and thyroid hormonal systems. EPA is using an initial Tier 1 screening battery consisting of five in vitro and six in vivo assays to evaluate a chemical’s potential to interact with the hormone systems in mammals and other animals. By order of the Office of Management and Budget, EPA must also consider Other Scientifically Relevant Information (OSRI) that is directly or functionally equivalent to data gathered in Tier 1, in lieu of developing new test data. This study characterizes the types of OSRI submitted by recipients of the first 67 test orders issued by EPA and reviews EPA’s approach to acceptance of OSRI. It also assesses the impact of OSRI acceptance on reducing the number of animals used in screening this first round of chemicals. Companies submitted OSRI in lieu of some or all Tier 1 assays for 47 chemicals and sought waivers for 412 assays. EPA granted only 94 waivers, an overall acceptance rate of 23%; of these, 50 were for in vivo tests. For 20 of the 47 chemicals, EPA denied all OSRI and required the entire battery of assays to be performed. In most instances, the OSRI accepted was either identical to data that would have been generated by the Tier 1 test or indicated a positive response by the chemical in question. Although identified as potential sources of OSRI in EPA’s guidance to test order recipients, guideline studies for pesticide registration, such as the mammalian two-generation reproductive toxicity study and 90-day rodent or dog studies, were all consistently rejected by EPA as satisfying Tier 1 data requirements. The 50 in vivo waivers EPA granted saved about 2,800 animals; however, nearly 26,000 were killed in the Tier 1 assays EPA required. The study concludes with a discussion of the implications for future use of OSRI in the EDSP.

The concern did not stop at evaluating long-term annual means for comparing to chronic, health-protective air monitoring comparison values (AMCVs). For evaluation of acute exposures, the agency generally uses 1-h AMCVs, which are not designed to evaluate 24-h results. Thus, the development of a 24-h AMCV would allow the TCEQ to evaluate 24-h formaldehyde data for possible health concerns. Critical effect dose-response data for irritation (e.g., eyes, upper respiratory tract) from acute (~400 ppb) and chronic (~200 ppb) human studies suggest a narrow range for the lowest reported effect levels, indicating these irritant effects are primarily concentration dependent. The TCEQ conservatively used the same point of departure (POD) that its chronic noncancerous AMCV is based on (NAAEL of 70 ppb) because the exposure duration (8 h/d) is more similar to the 24-h duration of interest than the 2-4 h exposure durations for the acute studies. Because the 8 h/d exposure was repeated 5 d/wk for 10 yrs, and irritation appears to be primarily concentration dependent, an 8-24 h exposure duration adjustment was not be necessary. Dividing the POD of 70 ppb by an intrahuman uncertainty factor of 3 results in a proposed 24-h, health-protective AMCV of 23 ppb. The 24-h AMCV falls between TCEQ’s 1-h (41 ppb) and chronic (8.9 ppb) noncancerous AMCVs. To date, there has been only 1 exceedance in 1999 of the proposed formaldehyde 24-h AMCV when compared to Barnett Shale monitored data.
studies/endpoints led by academic investigators. Twelve extramural grantees were selected by NIEHS through an RFA-based initiative to participate in the overall study design and conduct disease-relevant investigations using tissues and animals from the core study. While the study is expected to contribute to our understanding of potential effects of BPA, it also has ramifications beyond this specific focus. Through CLARITY-BPA, NIEHS has established an unprecedented level of collaboration among extramural grantees and regulatory agencies. The CLARITY-BPA represents a potential new model for filling knowledge gaps, informing chemical risk assessment, and identifying new methods or endpoints for regulatory hazard assessments.

1571 How Consistent Are the Derived No-Effect Levels (DNELs) in the European REACH Legislation?
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The new European REACH regulation places more responsibility than hitherto on manufacturers and importers of chemicals ("industry") to provide safety information. An important part of the development of a REACH Chemical Safety Report is derivation of Derived No-Effect Levels (DNELs) which represent "the level of exposure above which humans should not be exposed". In order to study the consistency, we compared DNELs presented by industry at the website of the European Chemicals Agency (ECHA) with those derived by us in our interpretation of the REACH guidance (Chapter R.8: Characterisation of dose [concentration]-response for human health, http://echa.europa.eu/documents/10162/13632/information_requirements_R8_en.pdf). There are various DNELs, e.g. representing short-term, long-term, inhalation and dermal exposure, as well as workers and the whole population. We limited our study to "worker-DNELs long-term" for inhalation route as they resemble occupational exposure limits (OELs). We found 24 substances for which (1) such DNELs were given in the ECHA chemical database (http://echa.europa.eu/web/guest/information-on-chemicals/registered-substances) and (2) a scientific basis for OEL had been published by the Swedish Criteria Group within the last 15 years at the University Arbetech Hälsa (https://gupuea.uib.gu.se/handle/2077/3194/local--en). The results were startling, as the consensus recommendations for protecting workers with occupational exposure to Cr(VI) require the use of respirators and screen creams. There are only a few tools available for the scientific assessment of the consistency.

1572 Animal Use for Testing Involving Unrelieved Pain and Distress.
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Each facility in the United States that uses live animals for research, tests, experiments, or teaching must submit an annual report to the U.S. Department of Agriculture (USDA) that includes "the common names and the numbers of animals upon which experiments, teaching, research, surgery, or tests were conducted involving accompanying pain or distress to the animals and for which appropriate anesthetic, analgesic, or tranquilizing drugs were (or were not) used" (9 CFR, Chapter 1 Part 2, Section 2.36). In accordance with the definitions in §2132 of the Animal Welfare Act (7 U.S.C. 54), it is not necessary to report birds, rats of the genus Rattus and mice of the genus Mus bred for use in research, or fish, amphibians and livestock or poultry used in agricultural research. In the 2010 USDA annual report on animal usage, a total of 1,134,693 animals were reported, with 97,123 of those reported as experiencing unrelieved pain and distress. Based on an analysis of data by the National Toxicology Program Interagency Center for the Evaluation of Alternative Toxicological Methods (NICEATM), 95% (91,997/97,123) of the animals reported to the USDA as experiencing unrelieved pain and distress were used for testing. Of these animals, 57% (54,889/97,123) were used for vaccine testing, and 38% (37,108/97,123) were used for toxicity testing. Most of the animals used for toxicity testing were used for safety testing and drug efficacy testing. NICEATM is currently investigating and promoting alternative test methods to further reduce the number of animals used in painful procedures. Supported by ILS staff under NIEHS Contract N01-ES-35504.

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Alternatives for acute systemic toxicity testing are one of the highest priorities of ICCVAM and NICEATM. These are the most commonly performed product safety tests worldwide and are required by multiple U.S. Federal agencies. Acute toxicity testing can involve large numbers of animals and result in significant unrelied pain and distress to test animals. High quality reference data are needed to evaluate alternative toxicity tests that may reduce, refine (enhance animal well-being and lessen or avoid pain and distress), and replace the use of animals for acute dermal systemic toxicity testing. To identify appropriate reference data for the acute dermal systemic toxicity test, NICEATM collected and analyzed data for 1897 substances. Rabbits were used for 28% (526) of the studies, and rats were used for 72% (1371). Of the 1897 substances, 84% (1598/1897) had data for both male and female animals, and 98% (1561/1598) of those substances were in the same GHS dermal hazard category. For the 37 substances that showed a different dermal hazard category between the sexes, female values were more often in a higher hazard category (21 for females vs. 16 for males). Two hundred forty six studies reported day of death. Approximately two thirds of the deaths (67% of male deaths [513/761]; 63% of female deaths [463/733]) occurred by Day 2 after a 24-hour dermal treatment on Day 0. Eighty-five substances had sufficient data to calculate dermal dose–mortality slopes. Dose–mortality slopes did not vary by species, sex, or GHS hazard category. As expected, the dermal dose–mortality slopes were lower than acute oral dose–mortality slopes. These data were used to design a proposed sequential test for acute dermal systemic toxicity, the dermal up-down procedure, to reduce the number of animals tested for acute dermal hazard classification. Supported by ILS staff under NIEHS Contract N01-ES-35504.

1574 Regulatory Acceptance of the BG1Luc Estrogen Receptor Transactivation Test Method.
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NICEATM coordinates an international interlaboratory validation study of the BG1Luc estrogen receptor transactivation test method (BG1Luc ER TA, LumiCell®) developed by Xenobiotic Detection Systems, Inc. In 2010, the validation study completed its goal to evaluate the usefulness and limitations of the BG1Luc ER TA test method to screen for substances in vitro ER agonist or antagonist activity. The international validation study was sponsored by NICEATM, with participation from the European Centre for the Validation of Alternative Methods, and the Japanese Center for the Validation of Alternative Methods. In 2012, NICEATM–ICCVAM released a test method evaluation report on the usefulness and limitations of the BG1Luc ER TA test method. ICCVAM recommended the use of the BG1Luc ER TA as a screening test to identify substances with in vitro ER agonist or antagonist activity recommended that the BG1Luc ER TA test method could be considered as an alternative to the existing ER TA test guideline (EPA OCSPP 890.1300/OECD TG 455). All 15 ICCVAM member agencies, including the US Environmental Protection Agency, concurred with the ICCVAM recommendations. NICEATM sponsored the new method for evaluation by the Organisation for Economic Co-operation and Development (OECD), which approved the BG1Luc test method and added the BG1 agonist protocol to the existing Test Guideline 455. The BG1 antagonist method has been adopted as OECD Test Guideline 457. Acceptance of the BG1Luc ER TA test method by U.S. and international agencies is an example of increased cooperation and collaboration to support the international adoption of scientifically valid test methods. BG1 will protect people, animals, and the environment, while reducing, refining, and replacing animal use. Supported by ILS staff under NIEHS Contract N01-ES-35504.

1575 Quantitative Risk Assessment As the Basis for a Proposed NIOSH Recommended Exposure Limit for Hexavalent Chromium Compounds.
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Sponsor: D. Dankovic.

To update the National Institute for Occupational Safety and Health (NIOSH) recommendations for protecting workers with occupational exposure to Cr(VI) compounds, all aspects of occupational exposure to and control of hexavalent
chromium compounds (Cr(VI)); e.g., chromic acid, CAS No. 1333-82-0; sodium dichromate, CAS No. 7782-12-0) were evaluated including toxicology, risk assessment, analytical methods, and industrial hygiene practices. Derivation of a proposed Recommended Exposure Limit (REL) was one component of the updated risk management recommendations. The NIOSH proposed REL was derived based on the results of a quantitative risk assessment (QRA) of lung cancer incidence. The proposed REL was previously based on the quantitative limitation of the 1975 analytical method; at that time NIOSH recommended that occupational carcinogens be controlled to the lowest feasible concentration. Data from a cohort of Baltimore chromate production workers were selected for analysis due to the availability of extensive exposure assessment data, information about smoking histories, strong statistical power, and relative lack of confounding exposures. Excess lifetime risk at the REL of 1 μg Cr(VI)/m³ was estimated as 69% (95% confidence limits=3-12) lung cancer deaths per 1000 workers. Based on these results, NIOSH proposed a REL of 0.2 μg Cr(VI)/m³ 8-hour time-weighted average exposure during a 40-hour workweek. The proposed REL recommendation is 95% effective for a 45-year-old man with smoking histories.
Skin corrosion or irritation refers to the production of irreversible or reversible damage to the skin following the application of a test substance, respectively. An effective way of predicting test substance toxicity is to make use of testing strategies which incorporate a range of alternative test methods. For the determination of skin irritation and corrosion, hazard assessments for both endpoints could be conducted using in vitro test methods that have been regulatory accepted (OECD TG431 & TG439). In the present study, skin irritation and corrosion evaluations were performed on Reconstructed human Epidermis (RHE) models i.e. EpiSkin and SkinEthic RHE. In the case of skin corrosion, GHS guidelines differentiate between non corrosive (NC) to corrosive (C) substances with 3 subcategories: 1A, 1B and 1C. The current evaluation of the test method was performed on 81 test substances from a wide range of chemical for each subcategory class (38 NC, 31 Cat1B/1C and 12 Cat1A) enlarging the evidence base associated with this method. Using the EpiSkin test method, within-laboratory variability (>87%) was assessed in 3 runs. Therefore a sensitivity of 98% and overall accuracy of 89% (with an accuracy of 1A, 1B/1C, NC, of 79%) were obtained. The test method able to discriminate 1A, from 1B and 1C classes with the highest well-prediction rate for sub-categorize the substances in comparison with the others validated methods, was submitted to OECD for scientific review and adoption. Adoption of the EpiSkin seems sufficient to fill the gaps in terms of sub-categorisation predictions leading to a significant impact on the sub-group transport package labeling.

Application of Systems Biology to Identify Molecular Mechanisms and Biomarkers of Lead (Pb) Neurotoxicity: Implications in a Developmental Origin of Alzheimer’s Disease.

L. Freeman, Health Sciences, Purdue University, West Lafayette, IN.

The heavy metal lead (Pb) can induce a wide-range of adverse health effects depending on dose and duration of exposure. During development the nervous system is most sensitive to Pb toxicity with epidemiological studies linking neurological deficits at and below the previous CDC blood Pb level of concern. Although the toxicity of Pb is extensively studied, the underlying genetic, epigenetic, and molecular mechanisms of Pb neurotoxicity are not completely understood. Moreover, recent studies link developmental Pb exposure with latent effects that do not appear until late in life, indicating a developmental origin of adult neurodegenerative disorders. More specifically, the latent overexpression of hallmark genes and proteins in Alzheimer’s disease (AD) are reported in these studies. This session brings together a group of investigators that are actively applying systems biology (transcriptomics and epigenomics) and targeted approaches to define the mechanisms and identify biomarkers of both the developmental and late-life neurological alterations associated with a developmental Pb exposure in a variety of model systems and in human populations. Topics cover a study with the zebrafish model on the genetic mechanisms of developmental Pb neurotoxicity with an emphasis on transcriptomic alterations to a human comparative transcriptomic study in young adults aiming at establishing biomarkers between early-life Pb exposure and AD. The session also addresses the transcriptomic and epigenomic pathways of the developmental origin of Pb-induced neurodegenerative alterations with a specific focus on AD in rodent and primate models. Furthermore, the mechanism by which Pb increases the formation of amyloid β plaques in a transgenic mouse model is discussed. Overall, this session highlights the latest findings on the genetic, epigenetic, and molecular mechanisms of Pb neurotoxicity linking neurodevelopmental and later life impacts to further deduce the developmental origin of Pb-induced neurodegenerative disease with a specific focus on AD.

Genetic Mechanisms of Developmental Lead Neurotoxicity and Links to Adult Neurodegenerative Disease Pathogenesis.

L. Freeman, Health Sciences, Purdue University, West Lafayette, IN.

It is well established that lead (Pb) exposure is detrimental to neurological development, but the underlying mechanisms of Pb developmental neurotoxicity are not clearly elucidated. Furthermore recent evidence supports the hypothesis that a developmental Pb exposure results in later lifespan effects including pathological features characteristic of Alzheimer’s disease (AD) proposing a developmental origin of Pb-induced adult neurological disease. The genetic and epigenetic mechanisms underlying and linking the immediate adverse effects of the developmental Pb exposure and the lasting impacts of the developmental exposure throughout the lifespan are yet to be determined. In this study, global gene expression analysis using the zebrafish model demonstrated that developmental Pb exposure results in immediate altered expression of genes associated with axon guidance, neurogenesis, and neurodegeneration. Furthermore, gene expression alterations were correlated to functional changes in the form of altered protein levels and a delay in axonal growth. While being an established developmental model, characterization of the adult zebrafish is limited and is just beginning to be explored for application in neurodegenerative disease pathogenesis research. Studies utilizing zebrafish embryos show that many genes with direct roles in AD are highly conserved in terms of sequence and function supporting the application of the adult zebrafish brain as a model to investigate the role of a developmental Pb exposure in the pathogenesis of neurological disease. To further our understanding of these genes, proteome, transcriptome and epigenome analysis and gene expression characterization was conducted. This study demonstrates that a strong degree of conservation occurs in regards to the presence and orientation of specific functional domains and in expression patterns. This study provides a framework for assessment of the influence of a developmental Pb exposure on late life neurological disease pathogenesis.

Prenatal Lead Exposure and Biomarkers for Alzheimer’s Disease in Humans.

M. Mazumdar1, 2, J. Environmental Health, Harvard School of Public Health, Boston, MA; 2Neurology, Children’s Hospital Boston, Boston, MA. Sponsor: L. Freeman.

Animal studies suggest that early life lead (Pb) exposure influences gene expression and production of proteins associated with Alzheimer’s disease (AD). This presentation will discuss recent studies in humans implicating early-life Pb exposure in the pathogenesis of AD. Aggregation of β-amyloid (Aβ) is a hallmark of AD pathology and plasma concentrations of Aβ are biomarkers that potentially could predict the risk of AD prior to the clinical manifestation of dementia. To evaluate the association between early-life Pb exposure and AD risk, the mean plasma Aβ concentration was measured in young adults and compared among those that had high umbilical cord blood Pb concentrations and those with lower cord blood Pb concentrations. Expression of genes whose products affect Aβ production and deposition was evaluated and was inversely correlated with umbilical cord blood Pb concentrations. Gene network analysis suggested enrichment in gene sets involved in nerve growth and general cell development. These data suggest that prenatal Pb exposure may influence Aβ-related biological pathways that are implicated in AD. Challenges in designing studies that investigate the contribution of early-life environmental exposures to adult disease will also be reviewed in this presentation.

Do Epigenetic Pathways Initiate Late Onset Alzheimer’s Disease (LOAD)? Towards a New Paradigm.

N. H. Zawia, Biomedical and Pharmaceutical Sciences, University of Rhode Island, Kingston, RI.

Cognitive decline and many of the hallmark pathological features of Alzheimer’s disease (AD) such as amyloid plaques and tau tangles are present in normal aging individuals and pose a challenge for distinguishing AD from normal aging. The majority of AD cases occur in the elderly; however, it is still unresolved whether AD is a disease of old age or whether it has earlier beginnings. The development of early onset AD (EOAD) seems to be largely genetic; however, late-onset AD (LOAD) may be influenced by epigenetic factors acquired during early developmental stages. LOAD exhibits numerous non-Mendelian anomalies that suggest an epigenetic component in disease etiology. The sporadic nature of >90% of AD cases, the differential susceptibility and course of illness, as well as the late age onset of the disease suggest that epigenetic and environmental components play a role in the etiology of LOAD. In this presentation, the evidence derived from primate and rodents (genomic and epigenomic as well as biomarker-specific) that AD has a developmental origin and that early life exposure can reprogram gene expression in old age through epigenetic pathways rendering the brain more susceptible to neurodegenerative diseases will be provided.

CNS Homeostasis of β-Amyloid, Plaque Formation, and Lead Toxicity.

W. Zheng, Health Sciences, Purdue University, West Lafayette, IN.

The hallmark of Alzheimer’s disease (AD) pathology is aggregation of β-amyloid (Abeta or Aβ). Previous studies in this laboratory have established that the brain barrier systems are actively engaged in regulation of the homeostasis of Abeta in...
Brain extracellular milieu (Crossgrove et al., Exp Biol Med 2005;230:771) and that lead (Pb) exposure can alter the property of the Abeta transport protein (LRP1) leading to abnormal accumulation of Abeta in brain tissues (Behl et al., TAAP 2009;240:245; Neurotox 2010:31:524). In the current study, we used transgenic PDAPP mice which over-express amyloid precursor protein (APP) and exhibit amyloid plaques in brain tissues to investigate if in vivo Pb exposure shortened the onset of Amyloid plaques in brain and increased the plaque aggregation. Our data show that in vivo Pb exposure resulted in an increased deposition of amyloid plaques in these transgenic mouse brains. Mechanistic investigation revealed that Pb reduced Abeta clearance from the central nervous system by inhibiting LRP1 at brain barriers. Pb also directly participated in physiochemical reaction with Abeta plaques in the test tube. Moreover, Pb increased the concentrations of other metals (i.e., Fe, Zn) in amyloid plaques in the mouse brain by synchrotron X-ray fluorescent (XRF) quantitation. Within the plaques, Pb concentrations were found to be significantly correlated with those of Fe and Zn. Our data support a role of Pb in the formation of amyloid plaques; how these findings may relate to human AD etiology deserves further investigation (Supported in part by NIH/NIEHS ES008146 and ES017055).

**1586 Bone As a Target Tissue for Environmental Toxins.**

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Bone brings to mind the strong, but light, organ that provides the skeleton for vertebrates. However, bone is more than just inert, ossous tissue. The bone marrow is a multifunctional organ that supports not only ongoing bone remodeling, but also provides the microenvironmental niche for hematopoiesis and regulates whole body energy homeostasis and as such, represents a significant target for environmental toxicants. Critical cell types in the bone marrow include multipotent mesenchymal stromal cells (MSCs) and hematopoietic stem cells (HSCs). MSCs are the source of all blood cell lineages and the bone-resorbing osteoclasts. The interaction between osteoblasts and osteoclasts creates a balance of bone formation and resorption, which is essential for maintenance of bone quality. There also is essential crosstalk between the mesenchymal and hematopoietic compartments that supports lifelong blood cell generation. Lymphocyte development, in particular, requires stromal cell support/interaction. Understanding how environmental toxicants disturb the interplay of bone marrow compartments requires attention given the rapidly aging population who are already at risk for loss of bone quality and immune suppression. We will explore new data suggesting that bone is responsive to many environmental toxicants, which may perturb the delicate balance between bone marrow cell types. A series of presentations will define interactions within both the mesenchymal and hematopoietic compartments and how exposure to toxicants may impact bone biology. Presentations will move from a broad, multispecies analysis of the effects of persistent organic pollutants on bone to more focused analyses of effects of ethanol and metals (lead, organotins, and tungsten) on bone and the bone marrow microenvironment.

**1587 Bone Tissue As a Target for POPs Acting As EDCs (an Overview from Wild Animals to Humans).**

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Since World War II, there has been an increase in age-standardized incidence rates of osteoporotic bone fractures in industrialized countries, with the Nordic countries taking the lead. The reason for this increase is unknown but the idea that exposure to endocrine-disrupting chemicals (EDCs) could be involved has been put forward. The consequences of environmental contaminant-induced bone toxicity is evident across many species. Exposure to persistent organic pollutants (POPs) acting as EDCs, has been shown to negatively affect bone tissue (e.g., lead, osteoporotic phenotype) in laboratory models (monkeys, rats, mice, frogs, goat, sheep, bone cells), as well as wild-life (polar bears, seals, alligators and herring gulls). Important signaling pathways such as the estrogen receptor and aryl hydrocarbon receptor pathways have been suggested to contribute to the observed effects. Importantly, epidemiological studies on humans also support this hypothesis since they show a relationship between exposure to endocrine disrupting POPs and decreased bone mineral density or increased risk of bone fractures. The literature on this topic is, however, rather small and prospective studies on in utero or early EDC exposure and future osteoporotic bone fractures do not exist. The mechanisms behind the deleterious effects of EDCs on bone tissue also need to be a focus of study.

**1588 Lead Exposure and Skeletal Dysregulation: A Molecular Mechanism of Toxicity That Contributes to Osteoporosis and Other Bone Diseases.**

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Rationale and Scope: Lead exposure and osteoporosis represent two of the most widespread health issues affecting humans. Our recent investigations provide evidence that these entities are linked. We have also identified a molecular mechanism by which lead can adversely affect bone forming cells. Experimental Procedures: We used a small animal rodent model as well as isolated cell experiments to evaluate the effect of lead exposure. The animals were treated with lead in their drinking water. The cell studies were performed in culture media with added lead acetate. All treatments achieved lead levels relevant to human exposures. Bone mass parameters were measured with micro CT analysis, histology and Raman infrared spectrometry. Cell effects and signaling pathways in osteoblasts were measured with molecular methods for mRNA and protein.

Results: Our findings indicate that lead exposure induces an osteoporotic-like phenotype in the animal model. Bone volume, bone formation rates, bone quality and bone cell number were all significantly depressed in the treated animals. Lead also induced a strong adipogenic response in the marrow space. Data from exposure of cells to lead indicate that all parameters of osteoblastic bone formation were also markedly inhibited. Immunohistochemical assays as well as a molecular analysis of in vitro cell signaling pathways indicate that the effect of lead is, at least partially, due to a strong depression of the TGFbeta and Wnt pathways. The mechanism for this depression appears to be related to an up regulation of the bone formation inhibitor, sclerostin.

**1589 Role of Nadph Oxidases and Reactive Oxygen Species in Regulation of Bone Turnover and the Skeletal Toxicity of Alcohol.**

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Recent studies with genetically modified mice and dietary antioxidants have suggested an important role for superoxide derived from NADPH oxidase (NOX) enzymes and other reactive oxygen species (ROS) such as hydrogen peroxide in regulation of normal bone turnover during development and also in the responses of the skeleton to toxicants such as ethanol (EtOH) which generate excess ROS in bone tissue. We have shown that EtOH causes bone loss as a result of NOX generated ROS, reduced bone formation via impaired Wnt-b catenin signaling and increased RANKL-dependent osteoclastogenesis. Buffering of anti-oxidant capacity though administration of the glutathione precursor N-acetylcysteine, completely prevented EtOH-induced loss of bone mineral density, inhibiting both suppression of bone formation and increases in bone resorption. There was no apparent bone phenotype associated with genotype at age 3 mo., in mice lacking the NOX co-factor p47(phox) and thus with impaired capacity to generate excess superoxide in response to EtOH. However, these mice were only protected from alcohol-stimulated increases in bone resorption. This was accompanied by inhibition of EtOH-associated increases in osteoclastogenesis and induction of RANKL. In mice where hydrogen peroxide concentration in bone is reduced as a result of transgenic expression of catalase, whereas 6 wk old mice had increased trabecular bone, 3 mo. old mice had reduced trabecular bone and impaired osteoblastogenesis compared to wild type mice and were unprotected against ethanol actions. These data suggest that ROS signaling involving hydrogen peroxide is important in regulation of adult bone formation but that excess production of superoxide via NOX as a result of exposure to toxicants such as EtOH can stimulate RANKL-dependent increases in bone resorption. Supported in part by R01 AA018282 (M.J.R.).
Tungsten: Effects on Bone Marrow and Lymphocyte Development.
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Very little is known about the toxicity of tungsten. Recently, increased environmental tungsten levels were found near sites of pediatric leukemia clusters. These leukemias were predominantly of the preB lymphocyte subtype of acute lymphocytic leukemia. While no data link tungsten exposure to leukemogenesis, tungsten is known to accumulate within the bone, the site of B cell development. This important hematopoietic compartment is likely the site where leukemogenic events occur. We have explored the effects of tungsten exposure on developing B lymphocytes using in vitro and in vivo models. In vitro, B lymphocytes are more sensitive to tungsten-induced DNA damage and growth inhibition than the supporting mesenchymal stromal cells. In wild-type mice, we have shown that tungsten concentrations rapidly increase in the bone and reach a plateau after approximately 4 weeks of exposure. Removal of tungsten results in a slow release of tungsten from the bone with a much slower "off" rate. In addition, we see alterations in bone mineral content and density following tungsten exposure. Tungsten exposure alters B lymphopoiesis and suppresses osteogenesis. Surprisingly, female-derived BM-MSCs are significantly more sensitive to TBT exposure than BM-MSCs derived from male mice. Tungsten exposure induces adipogenesis and suppresses osteogenesis. In vivo TBT exposure results in decreased cortical bone and increased marrow adiposity without significantly altering bone resorption. B cell populations are altered in both the bone marrow and the spleen of TBT-treated mice. In primary BM-MSC cultures, TBT potently induces adipogenesis and suppresses osteogenesis. Osteoclast and osteoblast dysfunctions have also been observed. In vitro TBT exposure results in decreased cortical bone and increased marrow adiposity without significantly altering bone resorption. B cell populations are altered in both the bone marrow and the spleen of TBT-treated mice. In primary BM-MSC cultures, TBT potently induces adipogenesis and suppresses osteogenesis. In vivo TBT exposure results in decreased cortical bone and increased marrow adiposity without significantly altering bone resorption.

Nonmonotonic Dose-Response Curves and Endocrine-Disrupting Chemicals: Fact or Falderal?
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“All substances are poisons. It’s the dose that makes the poison.” (Paracelsus, 1493–1541) is a fundamental tenet in toxicology: the severity of a response to a toxicant increases proportionally to the dose. Furthermore, it is generally assumed that dose-response curves for noncancer effects display a threshold below which there is no effect. Currently these assumptions are being challenged by claims that endocrine-disrupting chemicals (EDCs) often display nonmonotonic dose-response curves at low, environmentally relevant exposure levels; levels below traditional NOAELs (“Current Chemical Testing Missing Low-Dose Effects of Endocrine-Disrupting Chemicals—Endocrinology Society, 2012.”). In addition, the US EPA’s Endocrine Disrupting Screening Program (EDSP program) has been severely criticized, sometimes unfairly. This symposium will review the state of the science on EDCs concerning the shape of the dose-response curve in the low-dose range and the prevalence of NMDRCs. Talks will initially discuss mechanisms of action, the biologically plausibility for NMDRCs, and their focus on the molecular pathways that disrupt the estrogen and androgen signaling pathways and the prevalence of in vitro and in vivo NMDRCs. Presentations will also discuss the shape of the dose-response curves from “case studies” of estrogenic chemicals. These “case studies” are robust, mutagenic, and capable of inducing DNA repair in mammalian cells. The shape of the dose response curve in the low dose region has been debated since the 1940s, originally focusing on linear no threshold (LNT) versus threshold responses for cancer and noncancer effects. Recently, it has been claimed that endocrine disrupters (EDCs), which act via high affinity, low capacity receptors, commonly induce adverse effects displaying NMDRCs at low doses. The shape of the dose response curve in the low dose region has been debated since the 1940s, originally focusing on linear no threshold (LNT) versus threshold responses for cancer and noncancer effects. Recently, it has been claimed that endocrine disrupters (EDCs), which act via high affinity, low capacity receptors, commonly induce adverse effects displaying NMDRCs at low doses. Effects that would be missed in standard EDC screening and mutagenic testing protocols.

Nonmonotonic Dose-Response Curves (NMDRCs) Are Common after Estrogen or Androgen Signaling Pathway Disruption—Fact or Falderal?
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The basic biology of steroid hormone (SH) synthesis and action is reviewed to provide a molecular pharmacology basis for typical monotonic and nonmonotonic dose-response observations. Examples illustrate the molecular pharmacology paradigm for observed in vitro and in vivo effects. As the level of biological organization increases (from cell-free binding to transcription to in vitro adverse effects), the opportunity for complex nonmonotonic dose-responses emerge. Full strong receptor agonists and antagonists in vitro provide the highest level of predictability whereas weak agonists and antagonists are affected by ambient ligand levels. Agonist or antagonist activity can be passive (competition for receptor binding and subsequent direct receptor activation or inactivation) or active (influenced by co-activator and/or co-repressor preference and/or availability which can be cell type and tissue specific). In vivo cytoxicity, cofactor squelching and substance metabolism can confound data interpretation. SH ligands can also have rapid nongenomic effects leading to receptor/co-regulator phosphorylation and nonmonotonic responses. Similar to SH receptors, inhibition of rate limiting steroidogenic enzymes (delivery of cholesterol to the inner mitochondrial membrane and P450s) are more predictive than enzymes whose activity can be reduced by 95% or more without altering the rate of SH production. The intact animal leads to an even higher order of biological complexity where interactions with the hypothalamic-pituitary gonadal axis can logically lead to nonmonotonic dose responses. Exogenous administered androgens and the maintenance of spermatogenesis will be used as an example. Low levels of testosterone have little effect, physiological levels reduce GnRH and LH production, leading to reduced intratesticular levels of testosterone and the consequent inhibition of spermatogenesis and supraphysiological levels restore intratesticular testosterone and maintain spermatogenesis. The basic biology of steroid hormone (SH) synthesis and action is reviewed to provide a molecular pharmacology basis for typical monotonic and nonmonotonic dose-response observations. Examples illustrate the molecular pharmacology paradigm for observed in vitro and in vivo effects. As the level of biological organization increases (from cell-free binding to transcription to in vitro adverse effects), the opportunity for complex nonmonotonic dose-responses emerge. Full strong receptor agonists and antagonists in vitro provide the highest level of predictability whereas weak agonists and antagonists are affected by ambient ligand levels. Agonist or antagonist activity can be passive (competition for receptor binding and subsequent direct receptor activation or inactivation) or active (influenced by co-activator and/or co-repressor preference and/or availability which can be cell type and tissue specific). In vivo cytoxicity, cofactor squelching and substance metabolism can confound data interpretation. SH ligands can also have rapid nongenomic effects leading to receptor/co-regulator phosphorylation and nonmonotonic responses. Similar to SH receptors, inhibition of rate limiting steroidogenic enzymes (delivery of cholesterol to the inner mitochondrial membrane and P450s) are more predictive than enzymes whose activity can be reduced by 95% or more without altering the rate of SH production. The intact animal leads to an even higher order of biological complexity where interactions with the hypothalamic-pituitary gonadal axis can logically lead to nonmonotonic dose responses. Exogenous administered androgens and the maintenance of spermatogenesis will be used as an example. Low levels of testosterone have little effect, physiological levels reduce GnRH and LH production, leading to reduced intratesticular levels of testosterone and the consequent inhibition of spermatogenesis and supraphysiological levels restore intratesticular testosterone and maintain spermatogenesis. The basic biology of steroid hormone (SH) synthesis and action is reviewed to provide a molecular pharmacology basis for typical monotonic and nonmonotonic dose-response observations. Examples illustrate the molecular pharmacology paradigm for observed in vitro and in vivo effects. As the level of biological organization increases (from cell-free binding to transcription to in vitro adverse effects), the opportunity for complex nonmonotonic dose-responses emerge. Full strong receptor agonists and antagonists in vitro provide the highest level of predictability whereas weak agonists and antagonists are affected by ambient ligand levels. Agonist or antagonist activity can be passive (competition for receptor binding and subsequent direct receptor activation or inactivation) or active (influenced by co-activator and/or co-repressor preference and/or availability which can be cell type and tissue specific).
One aspect of endocrine disruptors is their ability to cause effects at low doses. Just as tiny amounts of hormones can have large effects on physiological systems, tiny amounts of chemicals that mimic hormones can have similar large effects. A feature related to low dose effects is the non-monotonic dose-response behavior of EDCs. In the past, basic toxicology focused on the simple dichotomy of toxic versus nontoxic, which implies that all substances can be harmful at high doses while at some lower dose, no harm is done. However, we now know that EDCs can create physiologically relevant effects at low doses, and these effects can have a substantial impact on our health. These effects may be beneficial at certain doses and deleterious at others. This modern understanding of non-monotonic effects is critical to understanding the behavior of chemical agents and also how resulting health effects may differ based on various exposures.

One of the challenges in addressing endocrine active compounds is how we define “low dose.” Standard toxicological dose-response models, typically deterministic in nature, are based on a toxicological framework that is not sufficient to address the complex behaviors and effects of low doses. A different approach to understanding low dose effects is the use of mechanistic models based on the current understanding of the mechanisms of action of the chemicals of interest. The use of mechanistic models to understand the effects of endocrine active compounds can be used to predict their behavior and impact at doses where standard approaches are not sufficient.

Endocrine-disrupting chemicals (EDCs) have been shown to affect reproduction, development, and metabolism in humans and animals. These chemicals can act as endocrine disruptors, mimicking or blocking the action of natural hormones. The effects of EDCs can include changes in reproductive health, altered hormone levels, and disruptions in the normal physiological processes that regulate metabolism, growth, and development.

Low dose effects of EDCs can be important to consider because they may occur at environmentally relevant concentrations. Understanding the mechanisms of low dose effects is crucial for risk assessment and regulation of EDCs. Traditional toxicological approaches often fail to capture the complexity of low dose effects, and more mechanistic models are needed to accurately predict and assess the impacts of EDCs at low doses.

There are several challenges in studying low dose effects of EDCs. One of the biggest challenges is the lack of adequate sensitivity in existing assays. Many of the standard assays used for toxicological testing are not sensitive enough to detect low dose effects. Developing new and improved assays that can detect low dose effects of EDCs is crucial for accurate risk assessment.

Another challenge is the lack of a comprehensive understanding of the mechanisms of low dose effects. While some mechanisms have been identified, there is still a need for further research to fully understand the mechanisms of low dose effects of EDCs.

Despite these challenges, there is growing recognition of the importance of low dose effects of EDCs. As a result, there is increased interest in developing new methods and approaches to study low dose effects. This includes the use of in vitro and in vivo models, as well as computational modeling and systems biology approaches.

Overall, the study of low dose effects of EDCs is an active area of research that is critical for accurate risk assessment and regulation of these chemicals. Continued efforts are needed to develop better assays and models, and to gain a deeper understanding of the mechanisms of low dose effects.

SOT 2013 ANNUAL MEETING
S 1601 Enhanced Susceptibility of Fatty Livers to Drug Hepatotoxicity and Innate Immune Responses.
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Steatosis is a risk factor for enhanced liver injury during drug hepatotoxicity and sterile inflammation. However, the mechanisms of this aggravated liver damage can vary dependent on the degree of steatosis and the pathophysiology. Steatosis can trigger microcirculatory disturbances, mitochondrial dysfunction and cause inflammation, which are important contributing factors for the increased susceptibility to liver cell death. The molecular mechanisms by which fatty liver promotes liver injury in genetic versus diet-induced steatosis models will be discussed in detail using examples of drug overdose (acetaminophen), hepatic ischemia-reperfusion and obstructive cholestasis.

W 1605 Health Impacts from Climate Change through Atmospheric Systems: Recent Findings and Challenges.
M. Bell. Environmental Health, Yale University, New Haven, CT. Sponsor: A. Farraj.

Climate change is anticipated to exacerbate many existing human health burdens such as through an increase in the duration and intensity of heat waves and through accelerated formation of tropospheric ozone. However, many questions on the nature of these impacts remain. This presentation discusses case studies of research on how atmospheric systems (air pollution, weather) could be impacted by climate change, thereby affecting human health, and will explore the challenges of this type of research. The case studies involve linking air quality, meteorological, and climate modeling to estimate the health consequences from changes in ozone levels in 50 U.S. cities and the mortality impacts from changes in heat waves in Chicago, Illinois, U.S. Challenges and assumptions of approaches to estimate health impacts from a changing climate include uncertainty in the various modeling systems (e.g., air quality models, climate change models), the use of current day relationships for weather or pollution and health for future scenarios, the changing distribution of susceptible subpopulations (e.g., elderly), and adaptive measures (e.g., air conditioning).

S 1602 Increased Risk of Drug Toxicity due to Altered Pharmacokinetics in Nonalcoholic Steatohepatitis.
N. J. Cherrington. Pharmacology and Toxicology, University of Arizona, Tucson, AZ.

Many severe adverse drug reactions occur when a patient is unable to metabolize and eliminate the standard dose of a drug due to alterations in drug metabolizing enzymes or transporters. Sources for this inter-individual variability include genetic polymorphisms or environmental factors such as inflammatory diseases that directly or indirectly alter the function of the enzymes and transporters that determine the pharmacokinetics of the drug. This presentation will discuss obesity-related and other possible sources of inter-individual variation in the transcriptional regulation, post-translational modification, and sub-cellular localization of specific drug metabolizing enzymes and transporters, thereby altering the pharmacokinetics and toxicity of drugs in patients that are at greater risk of adverse drug reactions.

S 1603 Biomarkers and Mechanisms for Steatohepatitis.
C. McClain. Pharmacology and Toxicology, University of Louisville, Louisville, KY.

Steatohepatitis may be caused by toxicants (TASH), obesity (NASH), or alcohol (ASH) among other causes. Mechanistic similarities and differences appear to exist between different types of steatohepatitis. We compared serum adipocytokines, cytokinin 18 (CK18), and antioxidants in human subjects with ASH, NASH, and TASH. All forms of steatohepatitis were associated with reduced serum antioxidants and insulin resistance (although adipokine levels differed by etiology). ASH and TASH were characterized by higher levels of pro-inflammatory cytokines. Likewise, ASH and TASH were associated with increased hepatocellular necrosis/apoptosis ratios compared with NASH which had a similar ratio to healthy controls. Lastly, complementary studies in mice showed major interactions between diet (high fat) and environmental toxins. This presentation will describe these findings and highlight the similarities and differences of novel clinical biomarkers of TASH, NASH, and ASH.

W 1606 Environmental Factors and Cardiometabolic Disease: Signals in the Air.
S. Rajagopalan. Cardiovascular Medicine, Ohio State University, Columbus, OH. Sponsor: A. Farraj.

Cardiometabolic diseases represent a pandemic, with the World Health Organization (WHO) projecting that more than 2.3 billion people will be overweight/obese by 2015. Technology innovations, globalization with its free movement of food and services, seismic shifts in agrarian practices coupled with rural-urban migration, nutritional transition to freely available high-caloric diets have irreversibly altered energy expenditures during work and leisure. These and other factors are helping to foster the continued "epidemiological transition" occurring across the globe. Scientific effort over the last few decades has focused on components of urbanization such as inactivity and dietary factors. More recent observations provide additional links between chronic exposure to environmental factors in air/water and propensity to diseases. This issue is of importance given the extraordinary confluence of high levels of airborne and water pollutants in urbanized environments. Multiple studies in China, India and other rapidly urbanizing economies demonstrate a steep gradient in urban-rural prevalence. This presentation summarizes recent evidence on how air pollution may represent an under-recognized yet critical linkage between urbanization and the emergence of cardiometabolic diseases, with a focus on diabetes mellitus.

A. K. Farraj. EPHD, NHEERL, US EPA, Durham, NC.

Dramatic reductions in air pollution over the last three decades have largely been driven by the enactment of federal regulations (e.g., the Clean Air Act in the United States). Today, policymakers and air quality managers rely on cutting-edge science to reduce and control air pollution. Toxicology is at the forefront of this effort providing critical input on health effects of air pollution including dose-dependence, the role of constituents and size, mode of action, and relative toxicity of air pollutants. Despite these advances, serious adverse health effects including cardiopulmonary mortality are still measurable at ambient air levels to which millions of people are currently exposed. Risk assessment of these air sheds is likely to get further complicated in light of the uncertainty posed by several emerging issues that intersect air quality and health. Climate change is one such issue that may affect health via direct effects of weather (i.e., heat and precipitation) and indirectly through increasing concentrations in ground-level ozone and particulate matter, two key air pollutants linked to adverse health effects. The burgeoning increase in obesity and associated metabolic disorders, groups with exaggerated sensitivity to the adverse effects of air pollution, is likely to aggravate health outcomes. With the implementation of new fuel standards and increasing popularity of alternative fuels, it is unclear what impact these changes may have on health effects of traffic-related emissions. Finally, several methods of power generation, including modern coal technology, nuclear energy, and hydrofracking have recently captured public interest, yet their impacts on air quality are unknown. This workshop discusses the current state of the science including key toxicological findings and recent innovations as well as challenges in the study of these emerging issues. The workshop concludes with a prospective look at air pollution research with a discussion session that engages the audience in an effort to define data gaps and potential mitigation strategies.

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The environmental impacts of combustion-derived vehicular emissions are well-established. In an effort to reduce greenhouse gas (GHG) emissions, improve fuel economy, and expand the nation's renewable fuel sector, new standards detailing the minimum volume of renewable fuel, which includes cellulosic biofuel, biomass-based diesel, advanced biofuel and total renewable fuel, contained in transportation fuel sold in the United States were recently proposed. Although it is assumed that these alternative energy sources will provide a "greener" fuel option, it is still largely unknown how their combustion emissions will impact human health, particularly when combined with traditional sources such as gasoline or diesel. Each biofuel has a unique chemical makeup, which will impact emissions characteristics and potential toxicity. Thus, this presentation highlights the new 2012 standards for renewable fuel, addresses the complexity of emissions arising from the combustion of various biofuel types, and examines the potential human health implications of incorporating biofuels into the transportation fuel sold in the U.S. For this last point, many of the health effects of vehicular air pollution in the last 10 years have
focused on particulate matter (PM). Recent studies suggest that certain biodiesels contain less PM and may be less toxic. Recent history of the industry is discussed with respect to chemical composition and component analysis, and toxicological effects, including cardiovascular, pulmonary and carcinogenic endpoints, in several rodent models.

1608 Fracking, Coal, and Nuclear Energy: Impacts of Contemporary Methods of Power Generation on Air Quality and Remediation Efforts.

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The complexity of the world's power production needs requires that a number of approaches be used to generate and distribute power. Natural gas, coal, nuclear, and other sources are all in play. Each of these energy sources is subject to intense regulation targeted at mitigating potential air and other contaminations, recently including mitigation of carbon dioxide. The increase in demand for natural gas has led to new methods for harvesting and obtaining it, including the controversial techniques such as fracking. Carbon capture and sequestration offers a number of engineering challenges, and methods for capturing include potential approaches that may initiate additional environmental concerns. Of course nuclear energy, especially considering the recent events in Fukushima, has its own environmental implications. This presentation focuses on emerging considerations related to the impact of power generation on air quality. The emphasis will be on coal emissions (modern vs traditional), carbon capture and sequestration (CCS), and natural gas (including fracking). Some mention of nuclear concerns will be considered. The presentation includes recent toxicology data on the health effects of inhaled amines and their degradation products used in CCS, and data on emissions from coal after they are atmospherically transformed in the environment and on air contaminants associated with fracking.

1609 Predicting the Future: Getting Ahead of Problems—A Presentation and Discussion.


Despite dramatic reductions in air pollution since 1970, the impact of pollutants on health and the environment remains a concern to both regulators and the general public. Studies have not been able to show thresholds of effect for prominent “criteria” pollutants like particulate matter (PM) and ozone (O3) and much uncertainty remains with air toxic air pollutants that vary across sources. Millions of people are exposed to the criteria pollutants, but there are subpopulations which are at risk due to unusual exposure circumstances or underlying biologic susceptibility or increasing age. Science continues to chip away at these concerns through human population studies, and human and animal toxicological studies. But as we look to the future with increasing national demand for energy and global population and industrial growth, there is increasing pressure on resources and the atmospheric reservoir for pollutants. Climate change with its impacts on air pollution chemistry as well as regional weather is widely thought to be undermining the gains we’ve seen in reducing the national air pollution burdens. New technologies, fuels, and ambient pollutant profiles require systematic assessments and innovative tools if we are to dissect these many interlaced issues to ensure optimal strategies to protect human and environmental health. This presentation attempts to draw on what we know and speculate on what we need to know from our science to forecast both emerging issues and solutions—preferably preventative—that have been discussed in this workshop. The audience will be engaged to embellish and enhance this discourse.

1607 The Current Regulatory Guidance on Neurotoxicity Assessment in Routine Preclinical Safety Studies is Inadequate and Needs Improvement.

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Neurotoxicity in routine preclinical safety studies is traditionally assessed with three H&E-stained brain sections and may be suboptimal. Hence, a recent study biodiesel-assessment of neurotoxicity in routine preclinical safety studies has been recommended but there is no expert consensus on the feasibility of incorporating modern methods of assessment into routine preclinical safety studies. We will explore and comment on the adequacy of the traditional approach to neurotoxicity assessment in routine preclinical safety studies. We will also examine the current regulatory guidance on neurotoxicity assessment in routine preclinical safety studies and discuss the feasibility of changing the current approach. Examples of emerging methods, such as MRI-directed histology and detection of circulating biomarkers of CNS damage, along with strategies for incorporating these techniques into standard preclinical safety studies will also be discussed. Finally, we will attempt to build consensus on appropriate approaches for improving the sensitivity of neurotoxicity assessment in routine preclinical safety studies by fostering a discussion between the audience and panel members. This discussion will focus on the feasibility of employing the proposed new markers, endpoints, and approaches and potential issues with interpretation of results of these studies. In conclusion, we believe that by examining the adequacy of current approaches of neuropathology assessment, discussing possible improvements to regulatory guidance and presenting emerging approaches of neurotoxicity assessment that this workshop will allow for a much-needed dialogue on the need and feasibility of improving the current methods of neuropathology assessment in routine preclinical safety studies.

1611 Routine Neuropathology Analysis for Nonclinical General Toxicology Studies.

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Microscopic toxico-pathological evaluation in Good Laboratory Practice (GLP)-type nonclinical general toxicology studies is one component of a battery of assays performed to assess potential nervous system toxicity of new chemical entities. Standard neuropathological assessment in such studies varies by species and among different industries and regulatory bodies. A Society of Toxicologic Pathology (STP) Working Group has recommended the following neuropathological strategy as suitable for all GLP-type nonclinical general toxicology studies. Selected brain areas should be examined routinely; caudate/putamen; cerebellum; cortex; hippocampus/pallidum; hypothalamus; medulla oblongata; olfactory bulb (in rodents only); pons; and thalamus. In rodents, these regions can be assessed in 6-7 full coronal sections on 2 standard-size slides; in non-rodents, they may be evaluated unilaterally in 6-7 standard slides each bearing 1 coronal hemisection. Spinal cord (cervical, thoracic, and lumbar regions) and peripheral nerve (sciatic and/or tibial) should be viewed in longitudinal/oblique and transverse sections. Current neurohistology practices—infiltration fixation in 10% formalin, embedding in paraffin, and initial analysis only of H&E-stained sections—is acceptable for nonclinical general toxicology studies. These recommendations slightly expand current practice (i.e., analysis of 3-4 transverse brain sections in rodents) to improve sampling consistency using available technology. These recommendations affirm the importance of consistent routine neuropathological evaluation in GLP-type nonclinical general toxicology studies while acknowledging that histopathology is only one component of neural safety testing in such studies. Therefore, institutions should retain flexibility in devising the neuropathology portion of GLP-type nonclinical general toxicology studies as long as major nervous system regions are evaluated systematically.

1612 A Regulatory Perspective on the Current State of Neurotoxicological Assessments in Drug Development.

R. Mellon. FDA/CDER/OND/DDAAAP, Silver Spring, MD. Sponsor: C. Toscano.

Neurotoxicological assessment of drug products is currently accomplished in preclinical studies by evaluation of central nervous system (CNS) safety pharmacology studies and general toxicology studies, which include simple behavioral observations and histopathological evaluation of the brain. Histopathological assessments in routine toxicology studies have identified adverse findings not detected by behavioral observation and therefore generate critical safety data. The scientific community has raised concern that histopathology in routine toxicology studies lacks adequate granularity for such a complex organ as the brain and therefore may not be appropriate for initial evaluations nor do they adequately identify when detailed “second tier” neurotoxicology studies are necessary. For example, the standard sampling and staining approach used in general toxicology studies may not be sufficient to detect certain CNS lesions (e.g., “Olney” lesions associated with NMDA receptor antagonists). Adopting an expanded sampling routine, such as the recently proposed Society of Toxicologic Pathology recommendations for the sampling and assessment of the brain in routine safety into Tier 1 testing, would increase the sensitivity of these studies. As there is currently no official CDER guidance on the assessment of the neurotoxicity of drug candidates, many approaches have been taken to addressing and reviewing the neurotoxicity of drug candidates; examples of these approaches and the level of success of such approaches will be provided. A regulatory perspective of the limitations of the current standard approach and feasibility of including modern methods of neuropathology assessment will be discussed. Overall, incorporation of modern methods of neuropathology assessment and improving the documentation of procedures used into the final study reports would provide greater confidence in the quality of the assessment. Such changes have the...
potential to impact the safety assessment of a drug candidate prior to first-in-human clinical studies, throughout drug development and as part of post-marketing safety assessments.

1613 Utilization of MRI Imaging to Optimize the Selection of Brain Sections for Assessment by Classical Neurohistopathology.

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Sampling of 3 coronal brain sections for morphological assessment in routine preclinical toxicology studies is clearly insufficient to reliably characterize all potential neuropathological changes that may be caused by new drug candidates. It has been estimated that 60 coronal sections would be required to adequately insure that this can be accomplished. This is never done in routine safety studies. Using a panel of classical neurotoxicants, including kainic acid, trimethyltin, domoic acid and nitropropionic acid that target different brain areas and structures and that exhibit effects varying from the massive to the subtle, a method has been developed for using MRI imaging to complement routine preclinical toxicology studies by allowing for a focused targeting and selection of candidate sections for histology in the rat. Proof of concept involves in-vivo MRI scanning to create files that can be compared and registered with coronal sections of the same brain sample after it has been fixed and stained appropriately for traditional neuropathology. Validation involves the establishment of concordance between the loci of pathological manifestation in each of these imaging approaches. Issues of comparability with respect to sensitivity, temporal equivalence of damage detection and whether MRI signals always predict morphological damage will be discussed. Successive in vivo MRI scans also allow subjects to be used as their own controls and for establishing time of onset, development and repair of morphologic damage. The MRI information may be used to optimize time of sacrifice for maximal visualization of traditional neurohistopathology. Most importantly, this method, when qualified, could provide guidance on where in the brain 3 ‘smart’ sections should be taken for classical neuropathology in support of preclinical drug safety studies. Overall, we show that low resolution MRI imaging in the intact rat, may represent a powerful biomarker that can inform classical neurohistopathology assessment, but is not a substitute for it.

1614 Translational Safety Biomarker Assessment of Neurotoxicity.


Neurotoxic effects of drugs pose a significant hurdle in the drug development process. Traditionally, neurotoxicity is assessed pre-clinically by classical histopathology, which has limited translational value for clinical development. Recent advances in imaging, functional assessment and the identification of fluid-based biomarkers of neurotoxicity might allow for the nomination of novel translational safety in neurotoxicity. Ideally, fluid based biomarkers of neurotoxicity detected in either CSF or blood could be combined with histopathology, high-resolution structural MRI and PET imaging, and other methods to assess neurotoxicity. We have assessed protein-based biomarkers of necrosis and apoptosis in addition to glial fibrillary acid protein (GFAP), ubiquitin C-terminal hydrolase L1 (UCH-L1), a biomarker of cell body injury and member of the ubiquitin proteasome system, myelin basic protein (MBP), a demyelination marker, and microtubule-associated protein (MAP-2), a biomarker of dendritic injury, in pilot studies with known neurotoxicants, including kainate. We developed and employed sensitive and qualified ELISAs to assess this panel of biomarkers. In addition, this data will be supplemented by existing multiplexed solutions for cytokines to incorporate inflammatory components in addition to these brain injury biomarkers. Along with the ELISAs, we have developed and applied multiple-reaction monitoring (MRM)-based panels of brain biomarkers. These MRM-type assays have shown significant improvement in sensitivity and allow for the multiplexed assessment of hundreds of proteins for targeted discovery with a follow-up verification studies on validated assays. These multiplexed biomarker panels comprise markers of different pathways, ranging from inflammation to synaptic changes and have proven to be useful in identifying neurotoxicity.

1615 Drug Safety Assessment and Regulatory Landscape in Emerging Markets.

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With 85% of world’s population and rapid economic growth, the emerging market countries are up and coming, and are now actively courted by large global pharmaceutical industries. Within the next decade, Asia is expected to overtake Europe in pharmaceutical sales. For example, China is predicted to be the second largest pharmaceutical market after the United States by 2015 (Nature Reviews Drug Discovery, 2010). Many large pharmaceutical companies have increased their presence in emerging markets for research and discovery, and are seeking to market their products locally. Due to the short history of innovative pharmaceutical research and development, many of the emerging countries have limited experience of first hand review of new drugs applications, hence rely mostly on the review decisions taken elsewhere to grant the approval in local markets. However, regulatory agencies of the emerging countries are working diligently to catch up with the international standards and in the process establishing the regulations that address their country specific needs to promote innovation and bring drugs that are appropriately tested to demonstrate safety for their own population. The establishment and expansion of regulatory agencies functions is creating local job opportunities for people with specialized skills, to help with the review of various components of the new drug applications while at the same time presenting challenges never experienced before due to the nature of innovation. This has sometimes led to regulations that are not well defined and open to interpretation. The drug safety assessment and regulatory landscape in China, India, Korea, and Brazil will be introduced during the presentations. The similarity and difference of the regulatory environment in the emerging countries will be compared with the major ICH guidelines. The challenges and possible solutions for companies seeking regulatory approval in emerging markets will be discussed. This will also provide an open forum for regulators from emerging markets to exchange ideas on how to tackle unique situation they experience.

1616 Framing the Workshop Theme.

S. Goel. Supernu Pharmaceuticals, Inc., Rockville, MD.

Countries in emerging markets are gaining increased focus regarding research and development activities and regulatory filings to market drugs due to an increase in consumer demand and rapid growth of their economies. To this end, it is essential to understand the progress and the intricacies in the regulatory filings among emerging markets. The speakers of this workshop will address several common questions on regulatory filing in their respective countries, including 1) The overall structure of the regulatory agencies and divisions under which various classes of drugs applications are submitted for reviews, 2) Comparison of various milestones in regulatory review process, preIND meeting, IND, SPA, end of Phase 2 meeting, and NDA, 3) Discovery and nonclinical safety studies required to support various stages of drug development and general comparison with ICH guidelines, 4) Requirements for country of origin of data, 5) Expertise and composition of non-clinical reviewers, 6) State of GLP compliance in contract research organizations within their countries, 7) Language used in regulatory submissions, 8) Regulatory outcomes and issue resolution.

1617 Current Status of Preclinical Safety Assessment of Drugs in China.


In China, Ministry of Health had regulated new drug registration and evaluation from 1949 to 1998. Starting from 1998, State Food and Drug Administration (SFDA) took over this function. There are three laws in China regulating the drug evaluation and registration process. In addition, SFDA has published 17 technical guidance for safety assessment of chemical drugs or biologics and 12 for traditional Chinese herbal medicines. Currently, in China chemical drugs or biologics are divided into 6 different registration categories. They are (1) investigational drug never marketed in any country, including synthesis or semi-synthesis natural sources or by fermentation; optical isomer; new combination of products, (2) drug with changed administration route, not marketed both domestically and internationally, (3) drug marketed outside of China, but not in Chinese market, (4) raw materials and formulations of drugs with changed acid or alkaline radicals or metallic elements, but no changes of pharmacology, (5) drug with changed dosing form, but not administration route, and (6) raw materials and formulations of drugs fitting in Chinese national standard. Moreover, there are 11 registration categories for traditional Chinese herbal medicines. Preclinical safety assessments vary according to the registration categories of new drugs. During early development, investigator develops strategies to determine safety of investigational drug according to the registration category. In 2003, SFDA officially issued the "Good Laboratory Practice (GLP) for Non-clinical Laboratory Studies" and required all preclinical safety assessments of drug to be conducted in GLP accredited labs. China has developed its own unique drug evaluation system, which ensures safe and effective drug products for patients.
India is emerging as a major center for research and development of new chemical entities for therapeutic use. The Drug and Cosmetic Rules, 1945 and all its amendments set the foundation for manufacture and sale of drugs. The “Schedule Y” of the above rule specifies the studies that must be done in animals to establish safety and efficacy of new drugs prior to trials in humans. However, until recently India has been a center for manufacture of generic drugs. The Government of India, Ministry of Health and Family Welfare established the Office of Central Drugs Standard Control Organization (CDSCO) headed by the Drugs Controller General (India) [DCG(I)] to regulate drug development. The current and up to date information on CDSCO can be found at http://www.cdsco.nic.in. In March 2011, the DCG(I) formed 12 New Drug Advisory Committee (NDAC) based on broad categorization of use of drugs on various systems in the body, such as Reproductive & Urology, Cardiovascular & Renal, Ophthalmology, Vaccines, Dermatology and Allergy, etc. The NDACs consists of expert members to advise on the matters of review and regulatory approval of clinical trials & new drugs. The charter for NDACs specifies that the drugs under development should consider relevance to population, innovation, and unmet medical need in India. India is unique in that it officially recognizes the development and marketing of traditional medicines (Ayurvedic, Siddha, and Unani drugs) for therapeutic use. India also prohibits certain drugs for manufacture and sale that it considers detrimental to its populations health. In India, the regulatory compliance of non-clinical studies is monitored by National Good Laboratory Practices (GLP) Compliance Monitoring Authority in Department of Science and Technology of Government of India. The Indian GLPs are based on the OECD guidelines due to initial close working relations between OECD countries in audit and GLP certification of Indian CROs.

Korea is another major emerging center for research and development of new pharmaceuticals. The 2011 Pharmaceutical Affairs Law had changed the regulations for manufacture, application, review, and approval of drugs. Korea Food and Drug Administration (KFDA), similar to US FDA, is a sole regulatory agency governing manufacture, application, review, approval, and surveillance of drugs. Within KFDA, the Pharmaceutical Safety Bureau with the Drug Evaluation Department is responsible for establishment of safety control system for drugs, promotion/support of the pharmaceutical industry, establishment of standards for the production of drugs and quality management, approval screening for drugs, and collection and evaluation of information on the side effects of drugs. Biopharmaceutical & Herbal Medicine Bureau and Medical Device Safety Bureau have similar roles in the relevant areas. As the research and development institute under KFDA, NIFDS (National Institute of Food and Drug Safety Evaluation) has played the key role in comprehensive research that is necessary for food and drugs regulation in Korea. It has been instrumental in the development and evaluation of toxicological and pharmaceutical methods. Center for Drug Development Assistance in KFDA supports for new drug R&D and regulatory affairs. The nonclinical safety studies at various stages of development are required according to ICH guidelines. Korean GLPs are based on the OECD GLP principles as a member country of OECD and Korean CROs have been frequently monitored jointly by three GLP authorities: KFDA, Rural Development Administration (RDA), and National Institute of Environmental Research (NIER).

In Brazil, for registration purposes, the medicines are classified in different categories, such as synthetics, biologics, phytotherapeutics. Synthetic medicines are classified as new, generic and similar. Each category of medicine is regulated by specific resolution, and its registration is granted for five years. The registration must be renewed by resubmitting the drug for evaluation at the end of each five year period. The registration of biological products is regulated by ANVISA since 2002. Due to the recent expiration of several patents for innovative biotechnology products, the interest in the production and sale of copies of these products is growing in Brazil. The regulatory framework for registration of biological products, (including new molecules or copies), was established in Brazil in December 2010 with the publication of RDC No. 55/2010. For development of this technical regulation, the regulatory authorities, networked and guided by the dynamics of evolving science, regulations, and processes based on the experiences from the more established markets/regions versus the individual interpretation and application of guidelines in the emerging markets. The Question and Answer session would provide an opportunity to deliberate current thinking on the subject and summarize the thoughts.

Vertebrate development is replete with instances of tissue fusion in which two tissues – each with its own complement of mesenchymal and epithelial cells – meet, adhere and eventually fuse into a seamless mesenchyme and epithelium. Such fusion is critical for proper morphogenesis of multiple organs, with failure implicated in several developmental defects: cleft lip/palate; closure defects of the neural tube; omphalocele (body wall); coloboma (optic cup); and hypospadias (urethra). Some of these defects are context-dependent – e.g., instances of cleft palate caused by failure of the palatal shelves to elevate, or hypospadias concomitant with impaired growth of the genital tubercle. Nonetheless, there is mounting evidence for fusion-specific defects (e.g., Pallister-Killian syndrome and VACTERL association). Furthermore, a predictive model built from ToxCast Phase-1 data and ToxRefDB mapped cleft palate to Gene Ontology terms for diverse biological processes (balanced accuracies of 0.66-0.76): negative regulation of epithelial cell proliferation; epithelial-mesenchymal transitions (EMT); TGFβ2 production and type-II TGFβ receptor binding; and neural crest migration. We have thus constructed a cell-level computational model of tissue fusion that incorporates cellular adhesion and viscoelasticity with paracrine/juxtacrine control of proliferation, apoptosis, EMT and cell migration. The model includes three growing tissues – only two of which normally fuse – to explicitly model failures of both fusion competence and specificity. As a first demonstration, we model secondary palate fusion by tuning the relative contributions of apoptosis, EMT and cell migration to match dissolution of the midline epithelial seam. This flexible model is a platform for computational toxicology that can elucidate the varying interplay of such cell behaviors in the normal course of different tissue fusion events and their role in differential susceptibility to chemical disruption. (This work does not reflect EPA policy).

Adverse Outcomes Pathways (AOPs) are used to describe the steps of a chemical interacting with a biological system from the subcellular to population level of observation. We investigated the published literature to develop qualitative AOPs for oxidative phosphorylation by mitochondria. Literature searches were carried out to identify chemicals and potential mechanisms of inhibition for each of the complexes of the electron transport chain (ETC) and for ATP synthase. Compounds for which sufficient experimental data could be located were then used to build plausible mechanistic pathway maps.
A dataset of over 200 compounds covering experimental data in a number of relevant screens (measuring individual complex activity, cellular ATP levels, reactive oxygen species (ROS) generation, oxygen consumption, mitochondrial membrane potential and cytotoxicity) was constructed. Seven model compounds were identified as having sufficient data and plausible mechanistic evidence to construct AOPs. The compounds used were: Rotenone, Thienoyfilluoracetone, 3-Nitropropionic acid, Antimycin A, Myothiazol, and Cyanide for mapping the ETC pathway, and Oligomycin A for ATP synthase. Using these representative model compounds an AOP was mapped for each complex from the molecular initiating event (MIE) to effects at the cellular level. The AOPs for inhibition of complexes of the ETC were then combined to create a single pathway.

We report the construction of AOPs for oxidative phosphorylation in mitochondria and their speculative extrapolation to in vivo endpoints such as cardio toxicity, neurotoxicity and hepatotoxicity. The AOP maps are described in terms of the normal physiological pathway processes and potential intervention points, with reference to experimental in vitro assays which were used to support the elucidated steps. Not all compounds with mitochondrial effects may go on to display in vivo toxicity. An AOP view of such pathways is an useful construct in providing information in a form suitable for assessing outcomes relevant to risk assessment.

## 1624 Interactive Data Mining of Toxicogenomics and In Vivo Toxicity Databases Using Chemotypes to Improve Chemical-Disease Prediction Inferences and Mode-of-Action QSAR Models.

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The Comparative Toxicogenomics Database (CTD, http://ctdbase.org) aims to shed light on the connection between chemical exposures and human health outcomes by inferring relationships via integration of curated chemical-gene, disease-and chemical-disease data from the scientific literature, Independent of CTD, structural knowledge via chemotypes, features linking chemicals to phenotypes, has been developed based on in vivo studies from toxicity databases, including regulatory sources at the US EPA and the US FDA. A chemotype is defined as a chemical substructure annotated with atom/bond properties that carry biological activity information. We explored whether the chemical-disease links in CTD could contribute to the further development of chemotypes, and whether mode-of-action Quantitative Structure Activity Relationship (QSAR) models for toxicity endpoints built around chemotypes could be used to validate CTD disease inferences or further delineate phenotypic effects. We applied a set of previously developed MOA models and chemotypes, e.g., for cleft palate, to the data found in CTD. Many of the same chemotypes (representing attributes of glucocorticoids, retinoids, gonzoles, dioxanes, and phthalic esters) were found to be associated with cleft palate in CTD, either by direct curration or inferred via common interacting genes. The structural feature space enriched with cleft palate chemotypes was highly populated by the chemicals deemed associated with this condition in CTD. This work demonstrates the ability to leverage data mining of these adverse effects, mice were exposed by inhalation to gas metal arc-stainless steel (GMA-SS) welding fume at 40 mg/m3 for 3 hr/d for 5 d a week for 2 weeks followed by gene expression analysis of whole blood cells, aorta, and lung at 4 hr and 28 d post-exposure. New features in Ingenuity Pathway Analysis (IPA) were used to analyze the expression data. The analysis results establish the importance of interferons and the IRF-family of transcription factors, predicting that they are activated at 4 hours and maintained through 28 days in both blood and lung. The IPA Upstream Regulator Analysis also predicted involvement of several toxicants, such as nickel and ozone, which have been detected in welding aerosols thereby highlighting the utility of Upstream Regulator Analysis for toxicogenomics applications. In addition, as an important drug-discovery tool, IPA predicted several compounds that might be useful to ameliorate the inflammatory phenotype. One such compound was LY294002, a PI3K inhibitor that has already been shown to be efficacious in a mouse model of asthma. The new Mechanistic Network Analysis in IPA was used to computationally construct a plausible hypothesis that TRIM24 inactivation may lead to activation of IRF7 and STAT1 and other upstream regulators to drive the gene expression profile after the first exposure. Finally, Downstream Effects Analysis in IPA predicted large increases in proliferation, chemotaxis, and trafficking of immune cells in the lung after 4 hours. In summary, IPA has a unique new set of capabilities to enhance the mechanistic understanding of toxicological datasets and to provide support for new hypotheses that can be tested in the lab.

## 1626 Mechanistic Analysis of Welding Aerosol Toxicity Using Ingenuity Pathway Analysis (IPA).

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Welding involves occupational exposure to an aerosol containing gases and metal particulates that induce adverse physiological effects including inflammation, immunosuppression, and cardiovascular dysfunction. To develop a deeper mechanistic understanding of these adverse effects, mice were exposed by inhalation to welding deposition of welding aerosol (WDA) for 3 hr/d for 5 d a week for 2 weeks followed by gene expression analysis of whole blood cells, aorta, and lung at 4 hr and 28 d post-exposure. New features in Ingenuity Pathway Analysis (IPA) were used to analyze the expression data. The analysis results establish the importance of interferons and the IRF-family of transcription factors, predicting that they are activated at 4 hours and maintained through 28 days in both blood and lung. The IPA Upstream Regulator Analysis also predicted involvement of several toxicants, such as nickel and ozone, which have been detected in welding aerosols thereby highlighting the utility of Upstream Regulator Analysis for toxicogenomics applications. In addition, as an important drug-discovery tool, IPA predicted several compounds that might be useful to ameliorate the inflammatory phenotype. One such compound was LY294002, a PI3K inhibitor that has already been shown to be efficacious in a mouse model of asthma. The new Mechanistic Network Analysis in IPA was used to computationally construct a plausible hypothesis that TRIM24 inactivation may lead to activation of IRF7 and STAT1 and other upstream regulators to drive the gene expression profile after the first exposure. Finally, Downstream Effects Analysis in IPA predicted large increases in proliferation, chemotaxis, and trafficking of immune cells in the lung after 4 hours. In summary, IPA has a unique new set of capabilities to enhance the mechanistic understanding of toxicological datasets and to provide support for new hypotheses that can be tested in the lab.

## 1627 Insights into the TBX2 Role in Cellular Senescence in COPD: A Bayesian Structure Learning Approach.

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COPD ranks among the top leading causes of death worldwide. The transcriptional factor T box 2 (TBX2) plays an important role in the COPD molecular etiology. A senescence hypothesis for the etiology of COPD has emerged. TBX2 and related genes have anti-senescence activity. Their expressions are suppressed in COPD, but elevated in many cancers. The purpose of this study was to identify the directionality of regulatory relationships between TBX2 and other genes involved in cellular senescence. From a compendium of lung epithelium cell microarrays generated using the Robust Multi-Array Average procedure, a subset of genes of interest was used for structure learning. The Bayesian Network Structure Learning approach was then used to learn the regulatory relationships. The Bayesian Network Inference with Java Objects tool kit (Banjo) was used, in each instance, to generate the single best-scoring direct acyclic network, given the gene expression data. In order to focus the large search space, an initial network of genes that TBX2 interacts with including ANF, NKX2-5, NDRG1, CDKN1A, PML, E2F, CDKNA2, and CDKN2B was identified. Simulated annealing and Greedy Search were the network search algorithms used. The results predicted that TBX2 regulates both COL4A3 and PML. Reports in the literature confirm that TBX2 interacts with both COL4A3 and PML. Subsequently, the original known network was updated with these two interactions, and a new iteration of the learning procedure initiated.
As part of this new iteration, the compendium subset with expression data of genes of interest was updated to include some novel genes identified from recent studies. The results identified a new TBX2 interaction partner including WNT3, WNT6, WNT7B, BTNL11, IL12RB1, IL13, IL13RA1, IL27RA, CTNNNd1, E2F4, and FOXO3. The published literature, a functional map scoring relationships based on microarrays from over 15000 publications (HepaMmr), as well as entries in a bio-informatics database (Mmmr) were utilized to validate the results. Thus, the results affirm the importance of TBX2 in the COPD etiology.

Predicting the Metabolism of Drugs Using Computational Approaches

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An increasing number of computational tools are nowadays available, which are claimed to predict drug metabolism. These tools could most probably be used in an extended way to consider potential drug metabolites earlier in toxicity prediction of for example, genotoxicity or carcinogenicity. Therefore it’s important to understand the quality and limitations of metabolism prediction tools if applied in a broader context. However, to date, the software packages used in metabolism prediction have only been validated with relative small datasets containing limited number of compounds. Hence, they require a more comprehensive and exploratory evaluation.

Aim: Meteor (Lhasa Ltd., Leeds, UK), an available prediction tool, has been used to predict the metabolites of 325 drugs in therapeutic use. The results of the prediction were compared to the known human metabolites.

Methods: The WHO ATC index was used to extract drugs for the therapeutic classes of antiinfectives (A07E, M01A, S01BA), central nervous system drugs and muscle relaxants (N, M03), cardiovascular drugs (C), oncologic substances (L01) and vitamins (J05, D06BB, S01AD). Additional information was received by queries in the DrugBank database. PharmaPendium and DrugBank were used to obtain information on the human metabolites of the therapeutic drugs observed in clinical studies. Meteor (version 13.0.0, Lhasa Ltd.) is a so-called knowledge-based expert system. As such it uses structured expert knowledge and its output offers the structures of metabolites with a guess on the probability of the occurrence of the metabolite. The evaluation shows that the reliability of the prediction of human metabolites is presently limited. Hence, further development is necessary to improve the prediction potential.

Comparison of Microarray and RNA-Seq for Gene Expression Analyses of Dose-Response Experiments

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Relative to microarrays, differential gene expression analysis using RNA-Seq data has been reported to offer higher precision estimates of transcript abundance, a greater dynamic range of transcript detection, and detection of novel transcripts. However, appropriate methods of analyzing RNA-Seq data and the number of reads required to match the sensitivity of microarrays are still a subject of debate. Moreover, comparisons of the two technologies have not uncovered clear cut-off levels for expression data.

Methods: The recent publication of the human genome sequence, as well as the recent release of genome sequences of many model organisms, make it becomes possible to comprehensively analyze gene expression in the context of gene function. Hence, we analyzed the expression changes of genes (orthologs) in the human genome.

Results: Our analysis indicates that RNA-Seq and microarrays both produce similar numbers of significant genes based on fold-change (FC ≥ 1.5) at p < 0.05. However, RNA-Seq produced significantly less genes based on p-value (FDR < 0.05) at even higher read depth of 9.3 M reads per sample. Among the significant genes, ~50% overlapped between RNA-Seq and microarrays at each dose. This result highlights the need for additional studies examining the use of RNA-Seq data in toxicology.

The Symbiosis of Mentoring: Getting the Most out of the Mentor-Mentee Relationship

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Mentoring is a critical element in the career development of all toxicologists, both in terms of making the most of potential mentors and developing effective mentoring skills. Whether through involvement in academia or helping to develop the expertise of an early-career scientist, most toxicologists will provide mentoring at some point in their career. The mentor role serves to transfer knowledge, give advice and provide support to a trainee or developing scientist, while the mentee is relied upon by the mentor to provide active participation and input into the relationship. According to national polls, as well as SOT-specific surveys, one of the resounding topics of interest for developing scientists is mentoring from the broad perspectives of choosing the right mentor, down to developing the skills to become a sound mentor. Therefore, this session was designed to complement existing mentor matching opportunities offered through SOT. Speaker presentations will focus on: (1) the fundamentals of mentoring, including the different situations and roles in which the mentor-mentee dynamic may be encountered; (2) an introduction to the most commonly used mentoring techniques; (3) identifying characteristics of a strong mentor-mentee relationship; and (4) mentoring towards the future of science and the challenge to overcome more complex scientific problems. Speaker perspectives will address the mentorship role within academia, government, and industry. Attendees of this session will learn to identify a healthy mentor-mentee relationship and understand the benefits to each member of the collaboration. Mentoring topics discussed in this session will be applicable to scientists at every career stage through highlighting the basics behind a strong, mutually-beneficial mentoring relationship.

Role of Systems Biology in Characterizing Risk of Developmental Origins of Disease

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Systems biology is the study of complex interactions of biological components, such as nucleic acids, proteins, chemical reactions, cells, and whole organisms, at multiple levels of organization. The National Research Council recommends implementing the use of systems biology approaches in the risk assessment process. While recent advances have been made to prioritize chemicals for further screening, to better understand mode of action, strengthen weight-of-evidence, and eventually replace traditional in vivo animal model data with in vitro and in silico methods, these data have not been systematically considered in mainstream approaches for risk assessment which also largely focuses on adult populations. The goal of this symposium is to consider emerging knowledge and information from systems biology to inform risk assessment and decision making in the arena of developmental origins of disease. Growing evidence suggests that chemical exposures in early life can have immediate and long-term effects on the structure and function of organ systems, leading to the development of disease later in life. Systems biology approaches can aid in understanding molecular, cellular, metabolic, and morphological changes resulting from such exposures. These approaches, which include analysis of complex data sets from multiple experimental sources, interdisciplinary ‘omics’ tools, and computer simulation modeling, are increasingly being applied to better understand developmental origins of disease. We will present current systems biology approaches on how to better identify developmental origins of disease and its implications to the risk assessment process through several case studies.

Predictive Models of Developmental Toxicity

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Environmental chemical exposure during development can lead to birth defects which can have lasting effects through adulthood. In vivo data from both rodent and non-rodent species gathered using standardized prenatal guidelines are used in risk assessments to determine developmental toxicity potential; however these studies may not indicate the underlying complex mechanisms which lead to developmental toxicity. We used a systems biology approach to develop predictive models
that link in vivo outcomes (including malformations, fetal weight reductions, and pre-natal loss) with potential mechanisms of toxicity. This approach uses in vivo data from guideline and literature studies and over 3 million in vitro data points from high-throughput and high-content screening techniques on chemical-biological interactions. Specifically, these assays are obtained from biochemical, cell-free and cell culture assays, along with embryonic stem cells and zebrafish embryos. The predictive models developed confirm known pathways, such as retinoic acid receptor and transforming growth factor, and suggest novel pathways such as G-protein-coupled receptors and inflammatory pathways to be important pathways involved in developmental toxicity of environmental chemicals. These models have the potential to aid in the prioritization of chemicals for further targeted toxicity testing and risk assessment, to generate hypotheses about mechanistic pathways leading to adverse developmental outcomes, and to reduce cost and increase throughput of chemical testing.

**1633 Multiple Organ-Omic Integration for HBCD Developmental Neurotoxicity Hazard Identification.**

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Exposure to environmental chemicals during critical windows of development has shown to disrupt adult neurobehavioral function; however, the mode-of-action is not always well understood. A systems biology approach was taken to identify toxicity pathways after exposure to the major brominated flame retardant mixture, hexabromocyclododecane (HBCD), along with the individual chemicals that make up the commercial mixture to better characterize developmental neurotoxicity, strengthen its overall weight-of-evidence, and support its hazard identification for risk assessment. Short-term disruptions to the developmental profile of genes, proteins, and metabolites in the hippocampus, liver, and systemic circulation were identified after exposure to the commercial mixture and individual stereoisomers. Different molecular pathway profiles generated in several organs suggest the necessity to assess the effects from the individual isomers as well as the mixture when evaluating HBCD and can be used as a model for other environmental mixtures for chemical risk assessment. This presentation will demonstrate the utility of an omics approach to capture information on various facets of brain development that can be altered after environmental exposures and to provide focus for further toxicity studies in determining possible modes-of-action including methods to prioritize individual chemicals in mixtures for chemical risk assessments.

**1634 Assessing Antiandrogenic Effects of In Utero Exposure to Dibutyl Phthalate and Other Xenobiotics.**

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The rates of male reproductive tract malformations such as cryptorchidism and hypospadia have increased in the United States throughout the latter half of the 20th century. These disursions are associated with various factors of in utero development that can be altered after environmental exposures and to provide focus for further toxicity studies in determining possible modes-of-action including methods to prioritize individual chemicals in mixtures for chemical risk assessments.

**1635 Mechanistic Pathways Underlying Low-Dose PFOA Effects on Mammary Gland Growth.**

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Perfluorooctanoic acid (PFOA) is a synthetic surfactant and, in areas of high contamination in the US, has recently been found to have a probable link to several human diseases. Prenatal exposure to PFOA causes delayed mammary gland growth in female mice. Exposure to low levels of PFOA throughout gestation results in persistent mammary gland abnormalities in female offspring that remain until late in life. Although activation of peroxisome proliferator activated receptor (PPARγ) is thought to mediate PFOA-induced liver toxicity, the role of this transcription factor in the mammary gland is unclear. Genome-wide microarray analysis, protein expression approaches, and morphological analysis were used to identify molecular mechanisms involved in delays following prenatal PFOA exposure. Our studies followed female mice exposed to environmentally relevant levels of PFOA in utero spanning neonatal time-points to late adolescence. Through analysis of whole mammary tissues and isolated cell types, it was found that PFOA altered pathways involved in epithelial-stromal interactions, including endocrine disruption, leaving the mammary gland more susceptible to the development of disease. The dosing levels of PFOA used in our studies produce body burdens in mice that are similar to those found in humans living in highly contaminated areas, therefore, these findings have implications related to the human risk assessment of PFOA.

**1636 Molecular Mechanisms Linking Maternal Diet-Induced Obesity to Offspring Metabolic Syndrome.**


Population studies have suggested that the maternal nutritional status during key stages of development is able to influence offspring risk of metabolic diseases. Recent focus has been on maternal over-nutrition and obesity in programming metabolic disorder in offspring. This stems from human data which has correlated maternal body-mass index (BMI), maternal weight gain during pregnancy, and gestational diabetes to offspring BMI and risk of metabolic disorders such as type 2 diabetes. A large number of studies have reported dysregulation of key organs involved in glucose homeostasis and feeding regulation in the offspring of obese mothers. We have utilized a mouse model in which dams are fed a western-styled diet (highly palatable, high-fat, high-sugar diet) during gestation and lactation and taken a whole systems approach to establish mechanisms through which metabolic disease in the offspring arises. Our studies incorporate analysis at the whole body, tissue and cellular level. Offspring of these dams are hyperphagic from a young age, leading to increased body weight and the development of type 2 diabetes later in life. Hyperinsulinemia is an early observation in these offspring compared to offspring in the control group fed regular chow and we have associated this with impaired insulin signaling protein expression, especially in adipose tissue. We have employed both candidate and genome-wide miRNA and microRNA array approaches to explore the mechanisms by which the programmed changes in insulin signaling protein expression occurs and to identify novel genes and regulatory elements that are associated with metabolic disorders in the offspring. These findings have important implications for the identification of markers of disease risk and identification of susceptible populations to adverse effects resulting from sub-optimal early life exposures.

**1637 Toxicopigenomics, Disease Susceptibility, and Implications for Risk Assessment.**

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In the past, classical toxicology has largely focused on the genotoxic effects of environmental toxicants and chemicals. Recent studies have clearly indicated that environmental chemicals, along with their genotoxic abilities, can cause epigenetic alterations that, in concert, may lead to the development of a number of pathological states. The field of toxicopigenomics has since emerged from the combination of epigenetics, which studies the methylation of DNA, histone modifications, and chromatin condensation, and toxicology. During the last few years, excellent progress has been made in detecting altered epigenomic profiles in response to various chemical insults. Future investigations, however, are needed to link chemical
Epigenetic Alterations in Response to Exposure to Environmental Toxicants.

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Epigenetics is the study of heritable changes in gene expression that are not associated with alterations in DNA sequence. Epigenetic mechanisms include DNA methylation, post-translational histone modifications, and nucleosomal positioning along DNA, and are important for maintenance of normal cellular homeostasis. Disruption of epigenetic mechanisms may compromise the balance of the cellular genome and lead to development of various pathological states, including cancer. Accumulating evidence clearly indicates that various environmental toxicants, independently of their genotoxic abilities, can cause epigenetic alterations. These changes in the cellular genome are exhibited as decreases in global DNA methylation, aberrant gene-specific DNA methylation, changes in histone modifications, as well as associated alterations in chromatin status. Importantly, epigenetic alterations can be detected long before the occurrence of genetic changes; and are stable, persisting after the initial exposure-related DNA damage has been repaired. These findings highlight the significance of epigenetic alterations in the mechanisms of toxicity and carcinogenicity.

Application of Cancer Toxicoepigenomics in Identifying High-Risk Populations.


The epigenome is dynamic and very susceptible to environmental changes. Toxicoepigenomic studies are conducted to explain the epigenetic effects of environmental exposures. Epigenetic programming occurs during development and reflects altered gene expression in disease states. This programming differs from genetic polymorphisms or mutations because genetic changes are reflected in all cells, whereas epigenetic changes are cell- and tissue-specific. Studies that involve the measurement of epigenetic changes that occur at the genome-wide level and their association with disease are called genome-wide association studies (GWAS). Epigenetic alterations respond quickly to environmental changes, and technologies are available to measure these changes. During the last 5 years, excellent progress has been made in the field of altered epigenome profiling in response to toxins and environmental. It is now possible to use high-throughput technologies to measure epigenetic patterns that are altered by toxin exposure. Different environmental exposures, including toxins, affect different components of the epigenetic machinery and alter the methylation and acetylation equilibrium. Epidemiologic studies help to identify the etiology of a disease, especially factors that contribute to disease development. In the case of cancer, such factors include toxic substances, radiation, infectious agents, specific dietary components, tobacco, alcohol, and environmental factors. It makes sense to identify epigenetic markers in normal and exposed populations. The advantages of identifying epigenetic biomarkers of exposure include improved exposure assessment, documentation of early alterations preceding cancer development, and identification of high-risk populations. Examples of interaction between toxic substances and cancer initiation and progression will be discussed.

Changes in Histone Tail Modifications and Gene Expression in PBMC from Subjects Exposed to Nickel and Arsenic.

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A major target of Nickel (Ni) ions in cells is the iron- and ascorbic acid-dependent dioxygenase enzymes. The family of histone H3 demethylases is an example of these enzymes. Since Fe binding is required for catalytic activity and Ni ions readily displace the Fe from all of these enzymes, Ni exposure results in inactive enzyme which increases global levels of the activating mark H3K4 trimethylation (H3K4me3) and the silencing mark H3K9 dimethylation (H3K9me2). Arsenic (As) induces oxidative stress in cells and will deplete reduced ascorbate which is required for the activity of these histone demethylases. In vitro exposure to Ni or As increases the levels of posttranslational modifications of histone tails. Here we show changes in global levels of histone modifications in peripheral blood mononuclear cells (PBMCs) of subjects with occupational exposure to Ni by inhalation in a Nickel refinery in China and subjects from Bangladesh exposed to As in their drinking water. Occupational exposure to Ni was associated with an increase in H3K4me3 and decrease in H3K9me2. A global increase in H3K9me2 and decrease in H3K9ac was found in subjects exposed to As. Additionally, exposure to As resulted in opposite changes in a number of histone modifications in males compared to females (H3K9Ac) or histone acetylation, (H3K9ac) altered gene expression changes in PBMC in a smaller subset of individuals exposed to Ni and As. Hierarchical clustering revealed that several thousand genes were similarly changed across subjects highly exposed to Ni, and there was similar clustering with low Ni exposure and with controls. With As the results were more complex because of the strong effect of gender. With As there were genes that changed in a gender-dependent manner and also with As exposure independent of gender. The results of these two studies suggest that exposure to Ni or As compounds, and possibly other carcinogenic metal compounds, can induce changes in global levels of posttranslational histone modifications and gene expression in PBMC.

Chromatin Remodeling As a Driver and Target for Lung Cancer.

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Epigenetic silencing of genes and microRNAs (miRs) via chromatin remodeling and cytosine methylation has emerged as a major mechanism in lung cancer etiology. The reversal of gene and miR silencing using pharmacological agents and potentially dietary supplements (e.g., omega-3 polyunsaturated fatty acids [n-3 PUFAs]) by affecting cytosine-DNA methyltransferases (DNMTs), histone methyltransferases (HMTs), or histone deacetylases (HDACs) offers new strategies for prevention and intervention for lung cancer. We have developed an in vitro model using hTERT/cdk4 immortalized human bronchial epithelial cell lines (HBECS) that identifies molecular changes driving transformation of preneoplastic lung epithelial cells during exposure to carcinogens found in mainstream and environmental tobacco smoke. Our studies provided a mechanistic link between increased DNMT1 protein, de novo methylation of genes, and reduced DNA repair capacity that together seemed causal for transformation. Our recent studies show that the expression of HBECS to carcinogens for 4 wks induces a persistent and irreversible, multifaceted dedifferentiation program characterized by the epithelial-to-mesenchymal transition (EMT). EMT induction was epigenetically driven, initially by chromatin remodeling involving H3K27me3 enrichment, with ensuing DNA methylation sustaining the silencing of miR-200b, -200c, -205 implicated in genome-wide profiling. These miRs reversed transformation. Genome-wide profiling has identified >500 genes with a promoter CpG island whose expression is reduced after only 4 wks of treatment. The polycomb repressive complex PRCC regulated many of these genes. While exposing promoter methylation after 8 wks of further exposure to the carcinogens is seen in a small subset of these genes, many of these genes show dense methylation in tumor-derived cell lines and primary tumors. These findings implicate chromatin remodeling mediated by HMTs whose expression increases during carcinogen exposure and in primary tumors, as important initiators of epigenetic gene silencing (Supported by R01 ES008801).

Molecular Basis for Dioxin-Induced Cardiovascular Disease.

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Interactions between genetic and environmental factors are responsible for several forms of human congenital cardiovascular malformations. Of these, valvular stenosis and hypoplastic left heart syndrome are the most common forms of birth defects.
in humans, occurring in eight newborns from every 1000 live births and constituting 25–30% of all cases of human cardiovascular malformation. These two congenital heart defects are the leading cause of neonatal and infant death and, underscoring the fetal origin of adult disease concept, are also a major cause of adult cardiac insufficiency, linking fetal and adult cardiovascular disease. The main known risk factors for these abnormalities in cardiac development are genetic inheritance and maternal exposure to hazardous chemicals, but their precise molecular etiology remains elusive. We have recently found that treatment of differentiating mouse ES cells with dioxin represses Nkx2.5 gene expression and other cardiac markers as a consequence of AHR activation. By following the expression trajectories of cardiomyocyte markers during ES cell differentiation we found that dioxin silenced the expression of Nkx2.5 and other cardiomyocyte-specific genes, including the genes coding for cardiac troponin-T and α- and β-myosin heavy chains, and inhibited the formation of beating cardiomyocytes, a characteristic phenotype of differentiating mouse ES cells. The key mediator of these effects was the aryl hydrocarbon receptor (AHR), epigenetically silenced in pluripotent ES cells and activated during differentiation. Our work with ES cells may, therefore, provide insights into the epigenetic mechanisms responsible for human cardiac malformations due to environmental (organochlorinated compounds) or genetic (Nkx2.5 mutations) causes. Nkx2.5 repression by dioxin-mediated AHR activation may help us identify the regulatory loops controlling the Nkx2.5 function that determines embryonic identity and progression of cardiac tissue differentiation, and how these functions are silenced by the activated AHR. (Supported by NIEHS grants 5R01ES006273 and 5P30ES060969).

**1644 Noncancer Disease Risk Promoted by Low Level Arsenic Exposures.**

A. Barczowsky, Environmental and Occupational Health, University of Pittsburgh, Pittsburgh, PA.

Exposure to arsenic through drinking water and diet increases risk for a number of cancers and non-cancer diseases. Cardiovascular diseases contribute to a large portion of non-cancer arsenic-promoted morbidity and mortality, and may occur at relatively low levels of arsenic exposure that were considered safe based on cancer risk. While epidemiological studies have established the disease risk to low level exposures, basic studies have lagged in demonstrating mechanism for disease. However, recent low level exposure studies in mice and cultured human cells suggest novel receptor mediated and reactive oxygen species-dependent modes of action for arsenic induced pathogenic vascular and tissue remodeling, as well as metabolic dysfunction. This talk will review the current state of understanding of arsenic-induced cardiovascular disease risk and potential modes of action.

**1645 Exposure Assessment Methods for Dietary Arsenic.**

L. Barraj and N. Tran, Exponent, Washington DC.

A brief overview of types of dietary exposure assessments and discusses the associated data requirements will be presented. Existing food consumption data sources that can be used for estimating dietary exposure to arsenic are reviewed and their strengths and limitations are discussed. Algorithms and models typically used are presented and their applicability to the various exposure scenarios (e.g., short term versus long term) are discussed.
1650 Experimental Exposure of Mice to House Dust Alters Gut Microbiome and Attenuates Allergic Pulmonary Responses.

N. Lukacs, Pathology, University of Michigan Medical School, Ann Arbor, MI. Sponsor: M. Williams.

Atopic asthma is more common in clean Western countries and its prevalence has increased greatly during the last century, along with improved hygiene and housing conditions. Exposure to a more diverse microbiome during childhood correlates with a lower risk of developing asthma during later life. Recent data has suggested that a diverse microbiome is associated with growing up with older siblings and pets. Dietary sampling from the environment can directly impact the gut microbiome. Interaction between intestinal microbiome and Gut Associated Lymphoid Tissue (GALT) not only directs local immune responses but also appears to affect the outcome of antigen-driven responses in the respiratory system. Using 2 different mouse models of allergic asthma (using the model-antigen ovalbumin – OVA – and the clinically relevant cockroach allergen – CRA), we set out to study pulmonary immune responses after dietary supplementation with household dust collected from homes with or without pets. Daily dosing of sieved dust administered to 6-8 week old BALB/c mice via oral gavage for 2 weeks was used for controlled exposure. Following the dust treatment, mice were sensitized and challenged with CRA, or alternatively received a transfer of OVA-specific DO11.10 T-cells and a subsequent OVA challenge. In the dust exposed animals a significant decrease in Th2 cytokine protein expression and total IgE levels was observed in pet dust-treated animals exposed to allergen. Moreover, these mice showed reduced mucous hypersecretion in the lung, as determined by PAS staining and RT-PCR for gob5. Interestingly, phylochip analysis of the cecal content of these mice revealed that the pet dust treatment altered the microbiome in favor of Lactobacillus species. Altogether these data demonstrate that short-term oral sampling of dust from homes with pets, but not without pets, significantly alters the gut microbiome and can subsequently attenuate pulmonary allergen-induced immune responses.

1651 Translating the Airway Transcriptome into Biomarkers of Tobacco-Related Lung Disease.

A. Spira. The Pulmonary Center, Boston University School of Medicine, Boston, MA. Sponsor: M. Williams.

While cigarette smoke exposure is associated with lung cancer, COPD and a number of chronic inflammatory diseases of the lungs, only a fraction of smokers develop these diseases and there are currently no effective tools for identifying those smokers at highest risk for disease. Furthermore, we lack biomarkers for measuring the host response to inhaled toxins like cigarette smoke. This talk will focus on applying whole-genome gene-expression studies to airway epithelial cells in order to gain insights into the physiological responses to tobacco smoke exposure. The goal of the presentation will be to explore how individual variability in the airway genomic response to tobacco smoke associates with risk of tobacco-related lung diseases and how this information can be leveraged to serve as clinically-relevant biomarkers of disease activity in both lung cancer and COPD. This talk will be of interest to the biomedical research community including those with expertise in genomics, molecular biology, and epidemiology related to tobacco exposure and pulmonary disease.

1652 Immune-Mediated Adverse Effects of Drugs and Environmental Agents.

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Many adverse effects of drugs and environmental agents are immune-mediated. Exposure to drugs is easier to determine than most environmental agents, and most drugs can cause idiosyncratic (i.e. specific to an individual) adverse reactions (IDRs) such as skin rash, liver injury, bone marrow injury, and autoimmune, most of which are immune-mediated. This provides a clear example of how different people can be affected by agents in different ways. Most drugs associated with a significant risk of IDRs form chemically reactive metabolites, which suggests that a major factor is the ability of the drug to form a hapten. However, this observation is also consistent with the danger hypothesis because reactive metabolites can also cause cell injury leading to the release of danger signals that promote an immune response. There are also environmental agents that form reactive intermediates and lead to an immune response, e.g. poison ivy and hair dyes can cause contact hypersensitivity. However, not all drugs that cause IDRs form reactive metabolites so there must be other possible mechanisms. Other apparent mechanisms involve epitope-presentation or epigenetic mechanisms. For example, an example of an environmental agent that binds to MHC leading to an immune response is beryllium, which binds to HLA-DRB1 that has a glutamic acid at AA 69, and very low exposures to beryllium can lead to severe chronic lung disease in patients with this genotype. Other agents such as vinyl chloride can lead to autoimmunity. Although IDRs are frequently referred to as dose independent, this is not true; drugs given at a dose of 10 mg/day are associated with a very low risk of IDRs. This may reflect a threshold in hapten density required to induce an immune response. This suggests that most environmental agents are unlikely to cause immune-mediated adverse reactions; however, beryllium is an exception, and this threshold may be significantly lower for agents that act more directly to modulate immune responses. Environmental agents such as benzene can also lead to immunosuppression. Funded by grants from CIHR.

1653 Targeted 'Omics Research in the Regulatory Environment.

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The US EPA's Office of Research and Development (ORD) has recently implemented the "Chemical Safety for Sustainability" (CSS) research program. Much of this program is dedicated to "facilitating faster, more efficient, more certain, and sustainable chemical assessments and management decisions". Achieving these goals will require high-throughput and multi-chemical assessments of environments, living intact organisms, and surrogate (in vitro) systems. Single chemical assessment strategies alone cannot support these requirements, and will be supplemented using "omic-based approaches" to exposure- and health-based assessments. This presentation will highlight opportunities to integrate cutting-edge 'omics approaches into existing regulatory-based research frameworks. Special attention will be given to biomarker-based 'omics tools that can be used to link across environmental, biological (in vivo), and in vitro sample environments.


C. G. Markgraf1 and M. Guha1. 1Drug Safety Evaluation, Bristol-Meyers Squibb Co., Mt Vernon, IN; 2DSS, Merck, Kenilworth, NJ.

Preclinical assessment of abuse liability potential for new drugs with central nervous system (CNS) activity has been a recent focus of both regulatory and industry interest. The standard animal models used for evaluation of abuse potential for these pharmaceutical candidates are self-administration and drug discrimination. These models have been well characterized in both rats and monkeys with known drugs of abuse. However, unique challenges encountered when evaluating drugs with novel mechanisms or with difficult pharmaceutical properties raise questions: How can reinforcing properties be evaluated if there is no acceptable IV formulation? What comparator drug is appropriate for a candidate compound with a novel CNS-active mechanism? In these situations, using a nonstandard model may offer an attractive alternative. The promise of such alternative animal models is balanced by the challenge in data interpretation and in incorporation of the data into a comprehensive risk assessment. This workshop will start with a discussion of the regulatory requirements and standard data collected for a preclinical abuse potential assessment, followed by presentations on conditioned place preference (CPP) and intracranial self stimulation (ICSS)—both nonstandard animal models of drug abuse potential in a regulatory setting. The scientific rationale for their use, how each compares to standard animal models, and the use of their data in risk assessment and in an abuse potential package for submission to health authorities will be discussed.


K. H. Horn. Drug Safety Evaluation, Bristol-Meyers Squibb, Mt Vernon, IN.

Preclinical data from abuse liability assessments are required to support the registration and scheduling of new drugs with central nervous system activity. Despite the availability of a standard set of well characterized animal models (self-administration and drug discrimination), there may be circumstances for compounds with...
novel mechanisms of action where the standard models are not suitable or cannot be utilized for the lack of suitable comparators for training or due formulation issues. In these cases, investigators may wish to turn to alternative approaches to understand the abuse liability potential for drugs under development. The purpose of this presentation is to highlight the complexity and challenges associated with designing an abuse liability package for drugs with novel or unknown mechanisms of action where the use of standard assays may not be appropriate or when pharmacological properties prevent the use of standard assays. The first part of the presentation will focus on the standard pre-clinical models of abuse liability (self-administration, drug discrimination, and withdrawal/dependence). Following this introduction, the presentation will discuss specific circumstances, situations, and/or challenges that investigators may encounter during drug development which may require consideration of unconventional approaches.

**1656 The Assessment of Abuse Liability: The Regulatory Perspective.**

S. N. Calderon-Gutkind. CDER, FDA, Silver Springs, MD. Sponsor: M. Guha.

The assessment of the abuse liability of a drug is part of the evaluation performed by FDA, according to the requirements of the Federal Food, Drug and Cosmetic Act. Issues concerned with the assessment of potential for abuse and with scheduling a drug under the U.S. Controlled Substances Act (CSA) are the responsibilities of the Office of the Center Director, Controlled Substance Staff. When a new drug has structural or pharmacological similarity to a known drug of abuse, sponsors are required to thoroughly characterize its abuse potential and submit study results for scientific review. A drug’s abuse potential is determined relative to a pharmacologically similar or other appropriate comparator drug. The abuse liability assessment is based upon comprehensive evaluation of chemistry, pharmacology (preclinical and clinical), pharmacokinetics, and pharmacodynamic profiles of the drug, and adverse events observed in clinical trials, as well as anticipated public health risks that may follow introduction of the drug on the market. This presentation will focus on the current guidance transition for industry on the assessment of the abuse potential of drugs, and on the particular challenges encountered when assessing the abuse potential of drugs with novel mechanisms of action or pharmacological profiles.

**1657 Conditioned Place Preference As a Tool for Assessing Abuse Liability.**

C. L. Cunningham. Behavioral Neuroscience, Oregon Health & Science University, Portland, OR. Sponsor: C. Markgraf.

The conditioned place preference (CPP) procedure is commonly used to assess the rewarding and aversive effects of drugs in laboratory animals. This procedure is also used to study the role of learning in the control of drug seeking and relapse as well as the brain mechanisms underlying drug-induced learning and reward. However, the CPP procedure has not been used systematically in the assessment of abuse potential. This presentation will focus on important issues that should be considered when using the CPP procedure for such assessments. The basic procedure and its parametric determinants will be briefly described, followed by a review of its advantages and limitations as a tool for assessing abuse potential. This review will focus on relevant theoretical, methodological, empirical and practical issues that arise when comparing the CPP procedure to the most commonly used tool for assessing abuse potential, i.e., drug self-administration. For example, consideration will be given to the use of these procedures for assessing “reinforcement” vs. “reward” and to the conceptual issue of whether these procedures simply offer different strategies for measuring a common biological process. Literature showing similarities and differences in drug effects between these procedures will be briefly reviewed along with examples showing use of a known drug or a drug of abuse in place of the other. Although a more systematic evaluation of its utility for predicting drug abuse liability in humans is needed, CPP might have significant value as an additional tool for assessing abuse potential in laboratory animals.

**1658 Use of Intracranial Self-Stimulation for Abuse Liability Assessment.**

T. J. Hudzik. 1Neuroscience Research, Abbott, Ludwigshafen, Germany; 2Institute of Pharmacology, Pavlov Medical University, St. Petersburg, Russian Federation. Sponsor: C. Markgraf.

All drugs entering the CNS may initially be viewed as having a potential risk for abuse. As there is no single test or assessment procedure likely to offer an ultimate characterization of abuse liability, preclinical abuse liability assessment is based on using several experimental approaches that focus on establishing pharmacological equivalency to known drugs of abuse, ability to induce physical dependence, and reinforcing effects. This report discusses intracranial self-stimulation (ICSS; brain stimulation reward) as a tool that could complement existing assessment strategies. Many drugs of abuse facilitate ICSS in laboratory animals prepared with electrodes implanted into brain “reward” areas. In addition to fairly robust predictive validity, ICSS has a number of important advantages in studying: i) drug-drug interactions; ii) quantitative comparisons among drugs; iii) complex drug formulations, especially those intended to be abuse-resistant; and iv) multiple, repeated drug exposures. This report limits a review of the ICSS literature to that pertinent to its usefulness as a procedure predictive of the abuse liability of drugs, and argues for the standardization of experimental protocols to enable a more final assessment of its usefulness in this regard.

**1659 Impact of Nonstandard Animal Models on Preclinical Study Design for Meeting Regulatory Requirements.**

M. Kallman. Discovery and Translational Services, Covance, Greenfield, IN.

Although most pharmaceutical companies have transitioned to provide non-clinical data to address regulatory requirements for scheduling decisions on new drugs these studies have typically been the traditional dependence, drug discrimination, and self-administration studies. Alternative models of conditioned place preference and intracranial self-administration have been available as scientific tools for sometime but they have only rarely been conducted to support the regulatory environment. The reason for the emphasis on the traditional models will be considered. In addition, a discussion of the advantages and potential applications of the non-standard models for scheduling decision making may provide insight into the utility of these models. The emphasis of the talk will be what the future environment might be and how we can develop scientific data packages using these non-standard models to meet the future decisions for scheduling.

**1660 The Placenta in Toxicology: Target or Travel Agent?**

C. J. Bowman1 and W. Foster2. 1Developmental & Reproductive Toxicology, Pfizer Inc., Groton, CT; 2Department of Obstetrics & Gynecology, McMaster University, Hamilton, ON, Canada.

The placenta is a fascinating organ in its dynamic form and function appearing as a transient and critical tissue in the reproductive cycle. It is a collaboration of maternal and zygotic cellular layers whose major function is a conduit between mother and fetus focused on the growth and viability of the next generation whilst preserving maternal well-being. The unique physiology of biodistribution and metabolism of the placenta also render it sensitive as a target of toxicity. The human placenta is a complicated organ and understanding the comparative physiology of nonclinical species can be a critical component for drug and chemical safety assessment; both as a potential target of toxicity and as a presumed barrier to undesirable xenobiotics. As in much of toxicology, it is important to appreciate different perspectives including clinical, epidemiological, basic research, industry, and regulatory. It is only when this information is integrated and applied that we can appreciate the value and importance of basic research and the views of regulatory and industrial scientists.

**1661 How Predictive Are Animal Studies in Detecting Fetal Risks in Humans?**

G. Koren1, 2. 1The Hospital for Sick Children, The University of Toronto, Toronto, ON, Canada; 2Molecular Toxicology, The University of Western Ontario, Toronto, ON, Canada. Sponsor: C. Bowman.

Every year scores of new molecules enter the market. Because half of all pregnancies are unplanned, fetal exposure to these drugs is inevitable. Unfortunately, due to wide variability and differences between human placenta in anatomy and physiology, making animal models of placental transfer difficult to interpret. There is an urgent need for non human models to predict human risk/safety. The placenta is the only human organ that can be ethically harvested and kept alive in vitro for 4-6 hr. By cannulating maternal and fetal vessels, the perfusion model can be used to study rates and extent of drug transfer across the human placenta, as well as placental functions such as oxygen consumption, hCG secretion etc. Using placental perfusion, we have established a model of predicting in vivo drug transfer, correlating it with results of in vitro fetal and neonatal sampling (Huson et al, Clin Pharmacol Ther, 2011). Implementation of this model can revolutionize the prediction of fetal exposure to drugs in vivo.
Environmental pollution is shown to affect reproductive functions in humans. In some situations exposures are unavoidable, such as traffic-derived air pollution in urban areas. The relationship between gestational exposure to air pollution and fetal outcomes is receiving progressively greater attention. Epidemiological findings correlate low birth weight, higher frequency of preterm birth, neonatal mortality and compromised specific reproductive endpoints such as infertility with air pollution. The limitations of epidemiological studies are associated with difficulties of identifying individual exposure levels as well as the presence and management of many co-founder factors such as nutritional and social status and smoking that challenges the establishment of causal relationships. To overcome these factors our group conducted experimental studies (murine model) using realistic concentrations of air pollutants, which “mimic” human exposures. Experimentation using real-world exposures to air pollutants provided corroboration of epidemiologic data and was used to identify the pathophysiological mechanisms involved. Changes in estrous cyclicity, fertility, number of ovarian follicles, embryonic implantation index, fetal development and placentation were some of the outcomes observed. In summary this session will provide an overview of the effects of environmental air pollution on reproductive functions; in particular, adverse effects on pregnancy outcome and placentation.

Triazole fungicides are known to cause changes in the placenta and they are known to cause late post implantation losses in prenatal studies in rats. Studies were initiated to investigate if there is a connection between these effects and the general mode of action of triazole fungicides, aromatase inhibition, causing a reduction of circulating estradiol. In prenatal studies, estradiol was co-administered with the triazole fungicide during rat prenatal development to test whether estrogen could prevent triazole late fetal resorptions. Co-treatment with estradiol reduced post implantation loss to 9%, compared to 48% without estradiol. Placental changes (labyrinth and tropospongium degeneration) were also less severe with estradiol treatment, indicating that these changes are secondary to reductions in estradiol. Thus, placental pathology may be the cause of late fetal resorptions. Expert pathological evaluation indicated that placental pathology could not support late fetal development; a quantitative evaluation suggested that degeneration of at least 70% of the labyrinth correlated well with late fetal resorption. As rat placenta is histologically and biochemically different from human placenta, while guinea-pig placentas are quite similar, we studied the effect of the triazole fungicide using guinea-pigs. In this model, prenatal triazole treatment did not result in an increase in the incidence of late fetal resorptions, despite a profound reduction in circulating estradiol levels. Similarly, no placental changes were noted. These differences can be explained by the fact that the rat placenta is unable to produce estradiol whereas the placenta is the main source of estradiol maintaining pregnancy following implantation in guinea-pigs and humans. Despite the triazole fungicide-induced decrease in total circulating estrogen production, the local concentration of estradiol in the guinea-pig placenta is sufficiently high to support normal maintenance and development in the fetal compartment. Therefore, the guinea-pig is a better model to assess prenatal toxicity of compounds inhibiting aromatase activity.
their homes, learn about historic and recent poisonings then solve a toxic mystery. The "Strange Case of Jennifer Strange" is based on the true story of the water intoxication death of a California mother. The activities will be demonstrated, and the pedagogical advantages of the 5E model for science education will be discussed.

1668 Inspector Tox Outreach Activity.
D. Hardee1 and A. Schatz2. 1Department of Pharmaceutical Sciences, St. John's University, Jamaica, NY; 2Merck & Co., Whitehouse Station, NJ.
Inspector Tox was developed by the Mid-Atlantic SOT (MASOT) chapter as a means of introducing the basic concepts of toxicology to school aged children. This program includes Inspector Tox an investigator of harmful or toxic events and helps guide the audience through determining if several "toxic scenes" set in the house, backyard, and at the beach are relatively safe or dangerous. The various scenes are devised to introduce a mix of potentially toxic compounds including chemicals (pesticides), natural toxins (bee sting, poison ivy), UV radiation (sunlight), social/behavioral toxicants (alcohol, cigarette smoking, and poor diet), and even includes components that the audience might not initially recognize as toxic, such as food allergies (peanuts) and dust.

1669 Silly Science and Other Activities for K–12 Outreach.
M. M. Bourgeois. EOH, University of South Florida, COPH, Tampa, FL.
Activities to explore dose, toxicity and therapeutic windows of drugs will be demonstrated and include describing dosages using familiar measurements (i.e. tablets, teaspoons, medicine cups, glasses, buckets and kiddie pools) that are toxic to show children the different amounts of specific household substances (water, salt, sugar, baby aspirin, etc.) necessary to do them harm.

1670 Exploring Toxicology: Designing Learning Goals and Evaluation Strategies for Outreach Activities.
E. Shanle and K. L. Blank. Molecular and Environmental Toxicology Center, University of Wisconsin Madison, Madison, WI.
Molecular & Environmental Toxicology Center (METC) graduate students at the University of Wisconsin-Madison participate in toxicology outreach events to promote a better understanding of toxicology to the local community. METC currently has 37 graduate students, and 15 students (41%) have participated in at least one educational outreach activity over the past year. Two activities graduate students use to teach about toxicology are Carnation Intoxication and Tox Land. Carnation Intoxication uses colored dye added to the water of white carnations to show that plants can take up toxins from their environment along with water and nutrients. Learning goals for this activity are age dependent and include a basic understanding of toxicology and the concepts of bioaccumulation and experimental design. Tox Land is a game that takes participants on a path through global events related to toxicology while also highlighting local daily activities that have positive or negative effects on people, animals, and the environment. The learning goals for Tox Land are that participants are informed about global/local toxicologically relevant events. Both Carnation Intoxication and Tox Land are evaluated using survey questions that reflect the learning goals to assess what participants learn. Evaluations are used to improve activities in order to better inform the public about toxicology.

1671 Resources for Toxicology K–12 Education Outreach: Updating the SOT K–12 Website.
The Society of Toxicology's Education K-12 Sub-Committee is updating the SOT website to increase support for K-12 education activities, share resources across the membership and with the public, and increase effectiveness in interacting with educators, parents and students. One main goal is to work with the Society membership to create the most useful resource website to help support excitement for toxicology through K-12 education outreach. Ideally, this website would be accessible to the public, easily searchable and contain hands-on activities, PowerPoint presentations and recommendations for increasing K-12 education outreach. Many SOT members are active in toxicology outreach, and many more would like to but don't know where to start. Working with Society members, the K-12 Sub-Committee will include leads to successful activities, including past Annual Meeting activities and ongoing Regional Chapter outreach activities. This resource website will serve as a tool for those interested in getting involved in K-12 education outreach, as well as for those interested in learning more about what toxicologists do.

P. Wexler. Toxicology and Environmental Health Information Program, National Library of Medicine, Bethesda, MD.
Cosmetic products represent an immense global industry. It has been estimated that the $60 billion cosmetics industry uses some 12,500 unique chemical ingredients in personal care products. These products and their chemical components are subject to varying degrees of regulation globally. In the US, the US FDA’s regulatory authority over cosmetics is relatively minimal and different from that of other products regulated by the Agency, such as drugs, biologics, and medical devices. Although the US FDA can inspect cosmetic manufacturing facilities, respond to complaints of adverse reactions, and conduct research on cosmetics and their ingredients to address safety concerns, the Food, Drug, and Cosmetic Act does not subject cosmetics to US FDA premarket approval. The regulation of cosmetics in the European Union is considerably more stringent. A Safe Cosmetics Act of 2011 was introduced in the US House of Representatives in June, 2011, with the aim of ensuring that all personal care products are safe. It would establish labeling requirements and a safety standard, and issue guidance prescribing good manufacturing practices for cosmetics and ingredients. People, largely though not exclusively women, are exposed to vast quantities of cosmetics over their lifetimes. Compared with the safety of most other products, cosmetics are commonly believed to present a minimal, if any, risk, and are overlooked in discussions of hazardous substances. Are the regulations currently in place adequate? What is the US government’s viewpoint? What is the stance of industry and advocacy groups on the cosmetics regulatory framework? How does the European Union handle issues involving the safety of cosmetics? Recent scientific issues surrounding production, safety, and testing of cosmetics could well influence the regulatory arena in years to come. These include 1) the increasing use of nanomaterials in cosmetic products and 2) computational and high-throughput alternative test methodologies. Both these topics will offer a backdrop to current and future regulatory approaches.

1673 Safety Evaluation of Nanoparticles and Skin.
N. A. Monteiro-Riviere. Anatomy and Physiology, Kansas State University, Manhattan, KS.
This presentation will review the physicochemical properties such as size, shape, coatings, vehicles and charge on skin penetration of nanoparticles (NP) that may enhance or prevent toxicity. There are properties of a chemical/NP that determines its propensity to cause dermotoxicity, which is the ability to penetrate skin and subsequently interact with biological components that could elicit a toxicological response. Recent advances of NP in consumer products such as cosmetics indicate safety concerns. Penetration of NP in skin is a controversial subject, partly because many factors that influence absorption were not studied and opinions of some investigators were generalized and based on limited studies of a few NP, many on large particles and not nanosize. Discrepancies may relate to differences in NP composition, surface chemistry, vehicles or solvents, techniques and methods of exposure, and analytical analysis, and duration of the experiment. This review will show how formulation, surface chemical alterations and mechanical stressing of skin and species can modulate the results.

1674 US EPA Computational Toxicology Predicting Cancer and Noncancer Outcomes for Cosmetics and Industrial Chemicals.
N. Kleinstreuer. National Center for Computational Toxicology, US EPA, Research Triangle Park, NC.
EPA’s Computational Toxicology Research Program (CompTox), part of the broader Chemical Safety for Sustainability Research Program, is at the forefront of a major transformation in the field of toxicology in how chemicals are evaluated for potential effects on human health and the environment. EPA is partnering with the cosmetics company L’Oreal, pharmaceutical companies, European regulatory authorities, and other U.S. government agencies to research ways CompTox can decrease the cost of testing large numbers of chemicals, reduce animal use, identify...
targeted testing strategies, and understand potential mechanisms of toxicity. Broad compound libraries may be screened and prioritized using rapid, automated high-throughput screening (HTS) assays with human gene and protein targets. Multiple HTS assay targets, e.g. chemokine expression in human primary skin cells, are highly relevant to cosmetics safety assessments. CompTox has already screened more than 1,000 chemicals in over 650 HTS assays in the ToxCast (toxicity forecaster) research program, including many key ingredients in personal care products, fragrances, and cosmetics. The L’Oréal partnership provides collaborative research funding and access to existing cosmetics ingredients safety data, which are compared to the ToxCast results to determine the utility of CompTox tools in the safety assessment of chemicals in cosmetics. CompTox research has also developed predictive models for cancer and non-cancer toxicity endpoints, with a focus on the molecular and cellular pathways that are the targets of chemical interactions. Using computational approaches, EPA is building decision support tools based on ToxCast in vitro HTS results to help prioritize cosmetics chemicals for further investigation, as well as applying and refining predictive models for a number of adverse health outcomes. This abstract does not necessarily represent EPA policy.

Therefore, complete banning of animal testing of cosmetics and their ingredients has a tremendous impact with far reaching consequences. This presentation will cover the safety assessment process of cosmetics and their ingredients and the changes that have been introduced by shifting from legislation to a regulation (including special considerations for CMRs, nanomaterials...). Also, the actual status of validated in vitro methods will be given and the prospective for the near future will be discussed.

Cosmetic products present on the European market must be safe for the consumer. In Europe, product safety is based on the safety of the ingredients (chemical structure, toxicological profile, exposure) and is the responsibility of the manufacturer, first importer or marketer. In July 2013, the EC Regulation n°2013/775/EU will replace in all Member States the well-known Cosmetic Directive 76/768/EEC with its important sixth and seventh amendments. Under this RECAST, a so-called responsible person gets a key function in the follow-up of the safety of the products under consideration. This is not an easy task as the safety of cosmetics and their ingredients has to be guaranteed by the use of in vitro methods, replacing experimental animal testing. In the EU, during the last years the 3R principle (Refinement, Reduction, Replacement) of Russell and Burch has been implemented in the specific legislation of several types of products, including chemicals, pesticides, pharmaceuticals, food additives, ... but the cosmetics legislation is the only one in the world allowing only replacement methods, which additionally need to be validated.

The two most important laws pertaining to cosmetics marketed in the United States are the Federal Food, Drug, and Cosmetic Act (FD&C Act) and the Fair Packaging and Labeling Act (FPLA). The presentation will cover the specific legal authorities granted to FDA under these two laws and the regulatory tools FDA uses to implement them. These regulatory tools include regulations, guidance, inspections, import monitoring, and other programs. The presentation will also cover other activities important to the oversight of cosmetic safety, such as FDA’s Voluntary Cosmetic Registration Program, the agency’s adverse event monitoring systems, outreach to consumers and health professionals, and more. The role of FDA’s research and other scientific efforts will be discussed. Examples of recent activities in areas important to cosmetic safety will be highlighted. Additional authorities that have been suggested for strengthening FDA’s monitoring of cosmetic safety and labeling will also be presented.

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Compliance costs are an important issue for cosmetics. The role of the industry has been under consideration. This is not an easy task as the safety of cosmetics and their ingredients has to be guaranteed by the use of in vitro methods, replacing experimental animal testing. In the EU, during the last years the 3R principle (Refinement, Reduction, Replacement) of Russell and Burch has been implemented in the specific legislation of several types of products, including chemicals, pesticides, pharmaceuticals, food additives, ... but the cosmetics legislation is the only one in the world allowing only replacement methods, which additionally need to be validated.
toxicology where Tox-21c could have a significant impact is developmental neurotoxicity (DNT). It is believed that cytototoxic concentrations, in the presence of concomitant biochemicals, contribute to the increasing incidence of neurodevelopmental disorders in children. However, due to lack of DNT studies only very few substances have been identified as developmental neurotoxins. This study aimed to develop an in vitro approach using metabolomics and gene expression for DNT assessment. A 3D rat primary neuronal organotypic model was exposed from day 7 up to 21 to suspected (developmental) neurotoxicants including pesticides (carbaryl, chlorpyrifos, lindane maneb), drugs (fluconazole, isotretinoin, phenytoin, terbutaline, valproic acid) and metals (CaCl2, PbCl2). Mass spectrometry based metabolomics measurements and quantitative measurement of gene expression in different cell types (neural precursor cells, neurons, and glial cells) were performed. Treatment with the different compounds significantly modified the expression of selected genes, related to the different stages of neuronal and/or glial cell development and maturation. Moreover, the mass spectrometry analysis showed differences in metabolite levels between control and treated cells. Further analysis of the altered metabolites could give mechanistic insight into the DNT of these compounds. This study demonstrates that gene expression and metabolomic analysis can be sensitive endpoints for DNT assessment. Funded by the Swedish Research Council and the FDA.

**1683** PCB 95-Induced Dendritic Growth in Primary Cultures of Rat Hippocampal Neurons Is Dependent on mTOR Activation.

G. W. Miller, H. Chen and P. L. Lein, Molecular Biosciences, University of California Davis, Davis, CA.

Despite being banned since the late 1970s, polychlorinated biphenyls (PCBs) remain persistent environmental toxicants that pose significant risk to the developing nervous system. We recently demonstrated that the non-dioxin-like PCB 95 alters neuronal connectivity by promoting dendritic arborization in cultured hippocampal neurons via ryanodine receptor (RyR)-mediated, transcriptionally driven mechanisms. PCB 95 sensitizes RyR and this interaction requires FKB12. Interestingly, FKB12 also regulates mammalian target of rapamycin (mTOR), and the mTOR signaling pathway enhances dendritic growth via increased protein synthesis. Based on these observations, we hypothesized that mTOR signaling contributes to the dendritic promoting activity of PCBs. To test this hypothesis, primary cell cultures dissociated from hippocampi of wildtype (Sprague Dawley) postnatal rats were plated at high density and exposed to PCB 95 (2.2,3,5,6-tetachlorobiphenyl, 20 pM – 2 nM) in the absence or presence of rapamycin. Exposure to PCB 95 from days 7 to 9 in vitro increased dendritic growth in pyramidal hippocampal neurons in a concentration-dependent manner. Simultaneous exposure to rapamycin at 20 nM, a concentration that inhibits mTOR activation, ameliorated the dendrite promoting activity of PCB 95. Ongoing studies are confirming that PCB 95 activates the mTOR signaling pathway implicated in dendritic growth, and determining which of these effects are RyR-dependent and which are mTOR-dependent. This is a novel mechanism by which low level exposures to non-dioxin-like PCBs cause developmental neurotoxicity, and perhaps dysfunction of other cell types, such as immune cells, whose function is heavily dependent on mTOR signaling. Supported by NIH grants R01 ES014901 and P42 ES04699.

**1684** Diazinon and Diazoxon Impair Glial-Neuronal Interactions and Production of Neuritogenic Proteins in Primary Astrocytes.

D. M. Pizzuto1, G. Giordano1 and L. G. Costa1, 2, 1Department of Environmental and Occupational Health Sciences, University of Washington Seattle, Seattle, WA; 2Department of Neuroscience, University of Parma Medical School, Parma, Italy.

Many organophosphorus insecticides (OPs) are increasingly recognized as developmental neurotoxins. However, the mechanisms by which they exert their neurotoxicity remain unclear. This project focuses on a widely-used OP, diazinon (DZ), and its active metabolite, diazoxon (DZO), and their potential to impair astrocytes’ ability to foster neurite outgrowth. We have previously shown that astrocytes play a major role in fostering neurite outgrowth in neurons; in this project, a 50% decrease in the longest neurite length was observed in primary hippocampal neurons incubated with primary astrocytes previously exposed to 10 μM DZ or DZO. We have also shown that this effect is largely mediated by increased oxidative stress in astrocytes that results from exposure to DZ or DZO. To further elucidate how these compounds adversely affect astrocytes and disrupt normal glial-neuronal interactions, levels of the pro-neuritogenic extracellular matrix matrix proteins fibronectin and laminin were evaluated by Western blot analysis and confocal microscopy. Fibronectin and laminin have been extensively shown to play a vital role in a variety of neurodevelopmental processes, including neurite outgrowth and differentiation. At the same concentrations that significantly inhibit neurite outgrowth, both DZ and DZO cause a 30–40 % decrease in fibronectin and laminin protein levels in astrocytes. These decreases are prevented by astrocyte pre-treatment with the antioxidants melatonin and N-t-butyl-alpha phenylnitrone (PBN), further supporting the role of oxidative stress in the mechanism of neurotoxicity. These findings were confirmed by confocal microscopy analysis of extracellular fibronectin expression, which showed a 40% decrease after exposure to DZ or DZO. These results suggest that DZ and DZO impair neuronal development by adversely affecting astrocyte production and secretion of important neuritogenic factors.
Interactions of environmental and genetic risk factors for Parkinson's disease (PD) are believed to modify disease onset and progression in a patient-specific manner. We sought to examine such interactions by assessing sensitivities of neural progenitor cells (NPCs) differentiated from human induced pluripotent stem cells (iPSCs) to PD-relevant environmental toxicants. These iPSCs were derived from a patient (SM) with biallelic PARK2 mutations and preclinical PD as assessed by DaTScan, and from healthy control subjects (CA and CB). Mn exposure resulted in significantly higher ROS generation in SM than CA NPCs, but no difference in cytotoxicity or mitochondrial fragmentation was observed. Moreover, we found that SM NPCs accumulate significantly less intracellular Mn than CA NPCs. These results indicate heightened susceptibility of SM NPCs to Mn given the higher rate of ROS generation and the comparable cytotoxicity and mitochondrial fragmentation in the presence of significantly less intracellular Mn. While the accumulation of Cu was not different between SM and CA NPCs, it induced significantly greater ROS generation, loss of mitochondrial function and cytotoxicity in SM than CA NPCs. These differences in Cu exposure-associated phenotypes were not observed in the primary fibroblasts used to generate the iPS lines. Given the role of PARK2 in mitochondrial integrity, heightened sensitivity of SM NPCs to manganese and copper may be due to suboptimal mitochondrial function in cells that lack functional PARK2. Our results support the hypothesis that genetic alterations may increase sensitivity to specific toxicants. We demonstrate here that iPS technology can detect patient-specific and cell type-specific differences in vulnerability to PD environmental risk factors. Support: NIH ES016931 and NS078289.
compounds and Colcemid as an aneugenic compound. Both, MMC and EMS exert a clastogenic activity which lead to a dose-dependent increase in mononucleus-positive binucleates only. In contrast treatment with colcemid resulted in an increase in micronuclei in both mononucleated and binucleated cells. The high level of CREST+ micronuclei in mononucleated cells suggest that polyplid cells which have undergone an incomplete endomitosis might be the case. Our results suggest that the micronuclei in mononucleated cells can be used to investigate the aneugenic activity of chemicals in a fast and easy way, and can be included in the CB assay with human lymphocytes.

1690 Induction of Nucleotide Excision Repair Is Diminished in Heterozygous p53 Knockout Mice following Chronic Dietary Exposure to Aflatoxin B1

J. E. Muldner1, R. Mehta1, G. S. Bondy1 and T. E. Massey1. 1Pharmacology and Toxicology, Queens University, Kingston, ON, Canada; 2Toxicology Research Division, Health Canada, Ottawa, ON, Canada.

Aflatoxin B1 (AFB1) is produced by Aspergillus species, molds that grow on grains, oilseeds and spices. AFB1, is biotransformed in vitro into a highly reactive metabolite that binds to DNA, forming DNA adducts that may induce cancer if not repaired. p53 is a tumour suppressor gene implicated in both AFB1 carcinogenesis and the regulation of DNA repair. Male heterozygous p53 knockout mice (p53+/-; B6.129- Tg53XtcpN5, Tacomic) and their wild type controls were exposed to 0.2 or 1.0 ppm AFB1, in AIN 954 semi-purified diet for 26 weeks. Nucleotide excision repair (NER) activity of lung and liver nuclear protein extracts was assessed with an in vitro assay, using addicted plasmid DNA as a substrate. In wild type mice, repair of AFB1-N7-Gua adducts was 124% and 96% greater than control in lung extracts from mice exposed to 0.2 ppm or 1.0 ppm AFB1, respectively, and 224% greater than control in liver extracts from mice exposed to 0.2 ppm AFB1 (p<0.05). In p53(+/-) mice, repair of AFB1-N7-Gua was only 45% greater than control in extracts from lungs of mice exposed to 0.2 ppm AFB1 (p<0.05), and no difference was observed in extracts from lungs of mice treated with 1.0 ppm AFB1, or extracts from livers of mice treated with either AFB1, concentration. When comparing AFB1, effects on NER activity after normalizing to untreated tissue, the induction of NER activity was significantly attenuated in p53(+/-) mice compared to wild type controls in both lung and liver and AFB1, dose had a significant effect in p53(+/-) mice. In conclusion, the increase in NER activity seen in wild type mice following chronic AFB1 exposure indicates a homeostatic response to DNA damage. This homeostatic response was diminished or lost in p53(+/-) mice, which is consistent with p53 having a key role in regulating NER. (Supported by CIHR Grant No. MOP-89098 and GRDP).

1691 XPC Haplotypes Alter Nucleotide Excision Repair and Levels of UV-Induced Genetic Damage.

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The XPC protein encoded by the polymorphic XPC gene is essential for initiating global genome nucleotide excision repair (NER). Over 90 single nucleotide polymorphisms (SNPs) have been reported in XPC, but only a few were evaluated as disease risk modifiers with conflicting results. SNPs do not exist as independent variants in the genome but as combinations forming specific haplotypes due to disease risk modifiers with conflicting results. SNPs do not exist as independent genetic structural alerts as well. Considering the advantages with respect to the amount of required sample and potential throughput capacity, HTFT is therefore thought to be a good screening tool for genotoxicity in the earlier drug screening.

The XPC haplotype encoded by the polymorphic XPC gene is essential for initiating global genome nucleotide excision repair (NER). Over 90 single nucleotide polymorphisms (SNPs) have been reported in XPC, but only a few were evaluated as disease risk modifiers with conflicting results. SNPs do not exist as independent variants in the genome but as combinations forming specific haplotypes due to linkage disequilibrium (LD) between them. The impact of XPC haplotypes on DNA repair capacity (DRC) is not known. Using bioinformatics, we recently determined the haplotype structure of XPC and found a correlation between XPC haplotypes and genetic damage in smokers. In this study we test the hypothesis that XPC haplotypes influence DNA damage by altering NER capacity through modifying transcription and/or translation. To test this hypothesis, we exposed human lymphoblastoid cell lines representing different XPC haplotypes to low UV dose to induce pyrimidine-6-4-pyrimidone photoproducts (6-4PPs) and cyclopyrimidine dimers (CPDs) formation. Levels of these adducts (reflecting DNA damage) and their rate of removal over time (representing DRC of the cells) were determined to evaluate the effect of XPC haplotypes on NER. We found that XPC haplotypes do not only influence NER, but that the haplotype influence is affected by the adduct being repaired. To provide a mechanistic explanation to our findings, we determined XPC mRNA and protein expression levels in different cell lines as a function of time after exposure. Our data indicate that XPC haplotypes significantly influence the rate of translation and transcription. Additional in-depth mechanistic studies examining the effects of haplotypes on XPC transcription, translation, and function are currently in progress to clarify the role of XPC haplotypes in disease risk. (Supported by F31-019053; T32-07454; P30-006676; RO1-CA093592; CA123208.)

1692 Evaluation of High-Throughput Mutation Test with Proprietary Pharmaceutical Candidates.


[Background] High-Throughput Fluctuation Test (HTFT) is a bacterial gene mutation assay in Salmonella typhimurium with the same endpoint as that of the Ames assay. HTFT requires only a few mg of a test sample for the assay, which makes it possible to incorporate the assay into an early drug developmental stage. It is reported that the results in HTFT have a high concordance with the Ames assay results for well-known positive and negative control compounds. In the present study, to assess the availability of the test to early drug screening, we tested our proprietary pharmaceutical candidates in HTFT.

[Method] HTFT was performed for 47 of our proprietary pharmaceutical candidate compounds in TA100 and TA98 with and without metabolic activation system prepared from rat liver (S9). Most of the compounds tested in this study were predicted as not having any obvious structural alerts for mutagenicity by a commercially available in silico system, DEREK for Windows. Microplates were used during the pre-incubation and incubation period, and mutation was detected with a ph indicator, which reflects the bacterial growth in medium. All the test compounds were also evaluated in the conventional Ames assay using five strains and the results were compared with HTFT results.

[Results and Discussion] The sensitivity (the proportion of positives in Ames assay correctly identified by HTFT), specificity (the proportion of negatives in Ames assay correctly identified by HTFT) and the concordance between the both assay results for a total of 47 test compounds were 88% (23/26), 86% (18/21) and 87% (41/47), respectively. In conclusion, the results of the present study indicate that HTFT is a reliable assay with a high sensitivity and specificity for detecting genotoxic compounds even among pharmaceutical candidates without obvious mutagenic structural alerts as well. Considering the advantages with respect to the amount of required sample and potential throughput capacity, HTFT is therefore thought to be a good screening tool for genotoxicity in the earlier drug screening.

1693 Establishment of the Comet Assay and Micronucleus Test Using Chimeric PXB-Mice® with Humanized Liver.

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Genotoxicity studies have been performed as in vitro screening tests to predict carcinogenesis and genetic disorders in humans. Notably, the comet assay and micronucleus test may be used to detect the in vivo genotoxicity of test compounds and their metabolites in rodents. However, metabolic activities differ between humans and rodents. Thus, we developed humanized chimeric PXB-mice®—whose liver retains human-type metabolic activities—by using albumin enhancer/promoter-driven urokinase plasminogen activator transgenic/severe combined immunodeficiency disease (uPA/SCID) recipient mice. Using the PXB-mice, we performed a comet assay and micronucleus test of N-ethyl-N-nitrosourea (ENU). At the SOT 2012 Annual Meeting, we presented that a dose-dependent increase in the number of positive cells was observed (i) in the comet assay using liver, (ii) in the micronucleus test using bone marrow, but not (iii) in the micronucleus test using liver, which is probably because of insufficient mitotic condition in the liver. In the present study, we established a new type of PXB-mice by using cDNA-uPA/SCID recipient mice, which would be expected to supply a higher mitotic condition for human hepatocytes. Cryopreserved human hepatocytes from a 2-year-old Hispanic male PXB-mice were treated orally with ENU for 4 weeks. In conclusion, we established comet assays and micronucleus tests using a new type of PXB-mice to predict human genotoxicity in vivo.
The skin is the main route of exposure of many chemicals and cosmetic ingredients; therefore, Cosmetics Europe (formerly COLIPA) has funded projects to establish and evaluate more realistic models for genotoxicity using 3D reconstructed skin (RS) issues. The aim is to use these to follow-up on positive results from the in vitro genotoxicity battery[1], which has been criticized for its low specificity. The RS model, EpiDermTM, was combined with the micrornuc (MN) assay and the resulting “RSMN” assay exhibited good intra- and inter-laboratory reproducibility[2], and correctly identified 3 coded chemicals as being either positive or negative[3]. A detailed protocol for the RSMN assay was published, together with a harmonized scoring atlas for micronuclei[4]. We have extended the number of coded chemicals to 29 as part of the pre-validation process. Eight of these were true positives, 11 were false positives and 10 were negatives. There was an excellent specificity (88%), demonstrating that the RSMN has a good potential to improve the specificity of in vitro genotoxicity assays as a whole. The 8 carcinogens with a suggested genotoxic mode of action, 5 were correctly predicted. While this indicates that the model shows decent sensitivity, the total number of true positives was considered too low to draw a final conclusion about the sensitivity of this assay. Therefore more coded compounds will be tested in a next project phase with a focus on genotoxic carcinogens. Overall, these data support the use of the 3D skin EpiDerm™ model for genotoxicity testing of dermally applied chemicals. [1] Phupler et al. 2010, Reg Pharm Tox; [2] Hu et al. Mutat Res. 2009, 673(2):92; [3] Aardema et al. 2010, Mutat Res. 2010, 701(2):123, [4] Dahl et al. Mutat Res. 2011, 720(1-2):12. Sponsored by CosEur and ECVAM

**1695 Reduction of Ataxia telangiectasia Associated Death Rates in ATM KO Mice with Yel002.**

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Ataxia Telangiectasia is a devastating disease that affects 1 in every 40,000–100,000 individuals worldwide. 1% of Americans carry a copy of a compromised Ataxia Telangiectasia Mutated (ATM) gene, an important signaling protein involved in both DNA repair and apoptosis. Having two copies of defective or null alleles leads to the disease, symptoms including motor defects (ataxia), extreme sensitivity to ionizing radiation, immunodeficiency, and predisposition to cancer. Lymphoma rates, in particular, occur at nearly 100 times the normal. Despite all of the defects, patients with AT are mentally normal, and many promising young high school and college graduates have their lives cut short by 22, the median age of death. The only treatments are palliative, focused on treating infections that result due to the compromised immune system.

The Schiestl Lab investigates the nature of DNA damage and its repair. During a yeast based high-throughput screen for agents that mitigate the damaging effects of radiation, the candidate drug Yel002 was identified. In later trials on healthy mice subjected to lethal radiation doses, Yel002 was found to significantly increase survival, even when administered a day after the insult. As radiation damage largely via strand breaks in DNA, it was thought to act by upregulating the DNA repair process. As AT patients symptoms result from DNA repair deficiency, we decided to treat ATM KO mice with the drug weekly in a long term study to cut death rates. Previous data on this same mouse strain showed that these mice usually die of cancer in the sterile facility by 50 weeks on average. Mouse treated with a weekly injection of 75mg/kg Yel002 show far lower death rates – over 80% are still alive at week 60, and survival looks promising as it continues.

**1696 Classification of Genotoxic Mode of Action in TK6 Cells via a High Content, Flow Cytometric In Vitro Micronucleus Assay.**


This laboratory has previously described a high content flow cytometric method for scoring micronuclei (In Vitro MicroFlow®) in CHO-K1 cells that is capable of discriminating aneugenic from clastogenic modes of action (MoA). It would be useful to capture MoA information when studying other cell lines, for instance the human cell line TK6. Previous reports suggested that the proportion of metaphase cells representing a single aneugenic signature can effectively discriminate between clastogenic and aneugenic MoA. We therefore set out to combine a flow cytometric micronucleus scoring method with a technique for enumerating metaphase cells and evaluate the multiplexed assay’s ability to differentiate clastogenic and aneugenic responses. In order to accomplish metaphase scoring, the fluorescent reagent anti-histone H3 (pS28) and anti-H3-P was incorporated into the flow cytometric micronucleus assay procedure. TK6 cells were treated in a continuous exposure design with each of ten reference clastogens and seven reference aneugens. At the time of harvest (24 to 27 hr after treatments were initiated), cells were processed according to In Vitro MicroFlow kit instructions; with the exception that anti-H3-P was incorporated into the sample processing. Data acquisition occurred for 4 mins per sample and provided approximately 2,000 metaphase cells per replicate. Each of the genotoxicants was observed to cause increased MN frequencies and relative survival values to decrease in a concentration-dependent manner. Whereas the proportion of metaphase cells to total cells (as well as the proportion of metaphase cells to G2/M cells) were decreased by each of the ten reference clastogens, they were elevated by each of the seven aneugens. These data indicate that automated and multiplexed micronuclei and metaphase cell-scoring of TK6 cells provide strong aneugenic versus clastogenic MoA signatures that effectively discriminate between these classes of genotoxic agents.

**1697 Qualification of 96-Well High-Throughput In Vitro Micronucleus and Comet Assays.**

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We recently qualified 96-well high throughput versions of the in vitro micronucleus (MN) assay in CHO cells, and the Comet assay in TK6 cells. The MN assay evaluated the ten reference compounds in OECD TG 487 using MicroFlow™ kits (Litron Laboratories; 10 independent trials), while the Comet assay evaluated six reference compounds using the Japanese Center for Validation of Alternative Methods (jaCVM) Comet validation protocol (version 6.2; four independent trials). All trials were performed, using 7-10 concentrations in duplicate cultures with and without S9, with a concomitant metabolic activation (S9), to assess inter- and intra-experimental variability, as well as sensitivity and specificity. Using an empirical analysis of the results, it was possible to reduce the criteria for a positive response for micronuclei and hypodiploidy to 2- and 6-fold concurrent vehicle control values, respectively, thereby increasing sensitivity without an appreciable loss of specificity. Mitomycin C and cytosine arabinoside were reproducibly positive without S9, as were benz(a)pyrene and cyclophosphamide with S9; all four compounds produced a clastogenic signature. Colchicine and vinblastine were positive with and/or without S9, and both produced a significant increase in micronuclei and hypodiploid cells, indicative of an aneugenic mechanism of action. In contrast, di(2-ethylhexyl)phthalate, nalidixic acid, pyrene and sodium chloride were all negative in all trials and at all dose levels with and without S9 except for one concentration of pyrene in one trial +S9. Reproducible positive results were observed in the Comet assay for the genotoxins 2-aminoanthracene +S9, and 9-aminoacridine, ethyl methanesulfonate and methyl methanesulfonate –S9, while reproducible negative results were observed for the nongenotoxins cycloheximide and triton-X –S9. These results support the utility of these high throughput assays for genotoxicity screening in general and are being employed in the US EPA ToxCast program.
In conclusion, the RSMN assay that utilizes the EpiDerm skin model is more specific in detecting genotoxic effects. Exposure to the genotoxin, Benzidine, was negative in EpiDerm but positive in 3-Triazinylpropional, and Ethyl Methanesulfonate. The genotoxicity was assessed using TK6 lymphoblast cell line cultured beneath EpiDerm tissues. Analysis of the complete set of chemicals resulted in 93.3% Sensitivity, 100% Specificity, and 95.5% Accuracy. In addition, a co-culture system that utilized a lymphoblast cell line TK6 cultured beneath the EpiDerm tissues was used as a target to expand the relevance of dermally applied compounds to systemic carcinogenicity. Micronucleus induction in TK6 cells was assessed after treatment of the EpiDerm with MitomycinC, β-Propiolactone, and Ethyl Methylene sulphonate. The TK6 cells, as well as the EpiDerm tissue itself, were found to be positive. However, exposure to the genotoxin, Benzidine, was negative in EpiDerm but positive in TK6 cells.

In conclusion, the RSMN assay that utilizes the EpiDerm skin model is more specific in identifying dermal human carcinogens than existing in vitro assays. The approach of using the human reconstructed skin together with a human lymphocyte target cell line appears to be a good method of detecting genotoxic effects for rapid identification of chemicals and expanding the relevance of dermally applied compounds to systemic carcinogenicity.

Identification of compounds with the potential to induce DNA damage is an important component of the chemical safety evaluation process. Micronuclei (MN), a frequently used endpoint for genotoxicity studies, are formed by clastogens inducingacentric chromosome fragments or aneugens disrupting anaphase during cell division. The traditional in vitro micronucleus (IVMN) test uses trained individuals to manually count MN, which is both labor intensive and time consuming. Here, we compare the performance of two automated high throughput methods for MN detection using CHO-k1 cells in 384-well format. Method 1 is a high-content MN assay using the Thermo Scientific Arrayscan VTI. Method 2 is the Intellicyt Corp high capacity flow (HCF)-based High Throughput Flow Cytometry (HTFC) Screening System® using the In Vitro MicroFlow® (24-well plates). This comparison was conducted in a round robin study. The compounds used included clastogens (etoposide, camptothecin, methyl methanesulphonate, bleomycin), aneugens (colchicine, griseofulvin, vincristine, geldanamycin), and non-genotoxic ones (Malic acid, sodium chloride). Both platforms correctly identified the known MN-inducing compounds and the 2 non-genotoxic compounds. Screening of a 384-well plate required ~3h for the ArrayScan and 30m for the HTFC; the latter platform was also able to distinguish aneugens from clastogens. These data show that the output of the IVMN assay might be increased dramatically to permit screening large chemical libraries on a robotic platform. Supported by NIEHS Interagency Agreement Y3-ES-7020-01.

Human susceptibility to environmental carcinogens is highly variable. Epidemiological studies have identified genes that confer resistance; however, these studies are limited due to small sample sizes. We use Saccharomyces cerevisiae (yeast) to profile eukaryotic genes that confer resistance to environmental carcinogens, such as the potent hepatocarcinogen, aflatoxin B1 (AFB1). A yeast collection (yeast) to profile eukaryotic genes that confer resistance to environmental carcinogens, such as the potent hepatocarcinogen, aflatoxin B1 (AFB1). A yeast collection

**1699 Reconstructed Skin Micronucleus (RSMN) Assay in Normal Human 3-Dimensional (NHu-3D) Skin Model.**

M. Klauser, Y. Kaluzhny, V. Karetsky and J. DeLuca. MatTek Corporation, Ashland, MA. Sponsor: P. Hayden.

Safety assessment of new products for human use requires genotoxicity testing. Current in vitro assays have low specificity resulting in a high rate of false positives that may be due to use of transformed cell lines, non-physiological exposures, and lack of normal metabolism. Furthermore, the 7th amendment to the Cosmetics Directive banned in vivo genotoxicity testing in 2009.

EpiDerm is a 3D normal human cell-based epidermal model that is highly reproducible, contains a skin-like barrier, and possesses biotransformation capabilities including CYP450, GST, and UDP enzymatic activity. The RSMN assay utilizes topical application of test materials in a similar fashion to actual human contact. Test materials are dosed 2 to 4 times every 24 h in the presence of Cytochalasin-B and are harvested from the EpiDerm within 24 to 48 h. Altogether, 50 materials (24 direct genotoxins, 14 non-genotoxins, and 12 genotoxicants that require metabolic activation) have been analyzed in the RSMN assay. Analysis of the complete set of chemicals resulted in 93.3% Sensitivity, 100% Specificity, and 95.5% Accuracy. In addition, a co-culture system that utilized a lymphoblast cell line TK6 cultured beneath the EpiDerm tissues was used as a target to expand the relevance of dermally applied compounds to systemic carcinogenicity. Micronucleus induction in TK6 cells was assessed after treatment of the EpiDerm with MitomycinC, β-Propiolactone, and Ethyl Methylene sulphonate. The TK6 cells, as well as the EpiDerm tissue itself, were found to be positive. However, exposure to the genotoxin, Benzidine, was negative in EpiDerm but positive in TK6 cells.

In conclusion, the RSMN assay that utilizes the EpiDerm skin model is more specific in identifying dermal human carcinogens than existing in vitro assays. The approach of using the human reconstructed skin together with a human lymphocyte target cell line appears to be a good method of detecting genotoxic effects for rapid identification of chemicals and expanding the relevance of dermally applied compounds to systemic carcinogenicity.

**1700 The Use of 3D Human Dermal Equivalents to Assess Genotoxicity of Textile Dyes.**

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Introduction: The use of 3D-human reconstructed skin seems to be promising for safety assessment of chemicals. Compared to traditional 2D in vitro systems, which basically represent one cell type, this model can provide organ-specific toxicity, mechanical/biochemical signaling and cell-cell communication. Humans are daily exposed to textile dyes through skin contact with colored garments. Studies have pointed that some textile dyes can be hazardous to human beings. Considering that dyes, under conditions of perspiration, can migrate from clothes and penetrate into human skin, identification of safe textile dyes, related to dermal exposure, is relevant to avoid human health problems. Objective: Genotoxicological assessments of human skin, identification of safe textile dyes, related to dermal exposure, is relevant to avoid human health problems. Objective: Genotoxic assessment of textile dyes.

**1701 Comparison of Two High-Throughput Methods for In Vitro Micronucleus Assay Assessment Comparison of Two High-Throughput Methods for In Vitro Micronucleus Assay.**

B. Goodwin1, J. Bennett2, Z. Liu1, S. Bryce2, K. Luu3, K. Witt4, C. P. Austin1, R. R. Tice and K. Xia5, 1NCGC, NCATS, NIH, Rockville, MD; 2Lirion Laboratories, Rochester, NY; 3InteliCyte Corporation, Albuquerque, NM; 4Division of the National Toxicology Program, NIEHS/NIH, Raleigh, NC.

In Vitro Micronucleus (IVMN) assays are used to screen chemicals for their potential to induce DNA damage in vitro human dermal fibroblasts, but a weak genotoxic response was observed. Conclusion: The difference might be related to the test systems used in each study (2D and 3D cell cultures). 3D-human reconstructed skin affords better cellular environment, which could provide more reliable data of the genotoxicity of chemical (e.g. textile dyes).
Ultra-violet (UV) irradiation is the most ubiquitous environmental inducer of skin cancer and melanoma. UV exposure creates DNA photoproducts primarily cyclobutane pyrimidine dimers (CPD) and 6-4 photoproducts (6-4 PP). These cause mutations and block DNA replication, unless repaired by nucleotide excision repair (NER). Thymine repeats (TTT) on G-rich strands of telomeric DNA are hotspots for CPD formation. Telomeres are DNA-shelterin protein complexes at ends of chromosomes, reported to lack a defined NER pathway for CPD repair. Diminished binding or loss of shelterin proteins TRF1 and TRF2 at telomeres cause telomere deprotection and end-to-end fusions directly leading to genomic instability. Also TRF2 binds to the key NER protein XPF-ERCC1 and recruits it to telomeres. Our working hypothesis is that telomeric photoproducts are repaired at lower rates compared to genomic lesions and that this repair inhibition involves interactions between XPF-ERCC1 with TRF2.

We measured photoproduct formation and repair in telomeres purified from human osteosarcoma(U2OS) cells exposed to UVC via a novel dot blot assay. We report a method for direct quantification of photoproducts formed in vivo and ensuing repair based on a standardized qPCR assay for DNA damage. Photoproducts were detected in genomic and telomeric DNA using specific antibodies following exposure of cells to 10 J/m2 UVC and after recovery times of 0 to 12 hours. CPDs decreased in genomic DNA after 12 hours but persisted in telomeric DNA. Southern blot assays using radio-labeled oligonucleotides that bind to telomeric deprotected sequences were used to quantify critical UV damage at telomeres and elucidate NER pathway regulation by shelterin. Diminished binding or loss of shelterin proteins TRF1 and TRF2 at telomeres causes telomere deprotection and end-to-end fusions directly leading to genomic instability. Also TRF2 binds to the key NER protein XPF-ERCC1 and recruits it to telomeres. Our working hypothesis is that telomeric photoproducts are repaired at slower rates compared to genomic lesions and that this repair inhibition involves interactions between XPF-ERCC1 with TRF2.

We measured photoproduct formation and repair in telomeres purified from human osteosarcoma(U2OS) cells exposed to UVC via a novel dot blot assay. We report a method for direct quantification of photoproducts formed in vivo and ensuing repair based on a standardized qPCR assay for DNA damage. Photoproducts were detected in genomic and telomeric DNA using specific antibodies following exposure of cells to 10 J/m2 UVC and after recovery times of 0 to 12 hours. CPDs decreased in genomic DNA after 12 hours but persisted in telomeric DNA. Southern blot assays using radio-labeled oligonucleotides that bind to telomeric DNA confirmed enrichment of telomeres in purified fractions. To investigate possible mechanisms for repair inhibition, we tested directly effect of TRF2 on NER protein XPF-ERCC1 via enzyme activity assays. We discovered that nucleolytic activity of purified XPF-ERCC1 on a stem-loop DNA substrate was completely abolished by TRF2. Our results provide the first direct biochemical evidence towards quantifying critical UV damage at telomeres and elucidating NER pathway regulation by shelterin.
Ethylene oxide (EO), a reactive industrial chemical, induced alveolar/bronchiolar adenomas and carcinomas in the lungs of B6C3F1 male mice at atmospheric concentrations of 50 and 100 ppm. The purpose of the current study was to characterize the mode of action (MoA) for lung tumors induced by EO. Male B6C3F1 mice were exposed by inhalation (6 hours/day, 5 consecutive days/week) to 0, 10, 50, 100, or 200 ppm (4 weeks) or 0, 100, or 200 ppm (8 or 12 weeks) and examined for incidence of micronuclei in the peripheral blood (MNT), DNA damage (Comet assay), histopathology of the lung, and characterization of DNA- and glutathione-adducts and lipid peroxidation in the tumor target and non-target tissues. In general, reactive oxygen species-related adducts (8-OfdG, CrotodG, M1dG) were only minimally affected, whereas alkylated DNA adducts (O⁶-HeDG, N₁-HeDA, N₆-HeDA, and N₇-HeG) were increased more robustly. There was a dose-dependent increase in glutathione adducts (HESG) in all tissues, although severe GSH depletion was not noted. There were no treatment-related changes in the MN-RET (reticulocyte) frequency or %RET. Comet analysis of the lung revealed a dose-dependent, statistically significant increase in DNA damage at 50 ppm and above. There was no treatment-related histopathology in the lung, although a slight decrease in the proportion of Ki-67 positive cells was observed at 4 and 8 weeks in the terminal bronchioles. In summary, these observations reveal a complex mechanism of biomarkers of exposure and effect following inhalation exposure to EO with a terminal bronchioles. In summary, these observations reveal a complex sequela of the carcinogenic process. Therefore, the shift in K-ras mutation spectrum observed in EO-induced mouse lung tumors may be due to effects on intracellular signaling pathways and selection of preexisting mutant cells, rather than direct mutagenesis of the K-ras gene by EO.
1712 Deciphering the Role of TRAIL in Altering Spermatogenesis by Controlling Germ Cell Apoptosis.

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TRAIL (TNFSF10/Apo2L) is a member of the tumor necrosis factor (TNF) superfamily of proteins, expressed in human and rodent testis. TRAIL is known to induce apoptosis via binding to its receptors DR4 (TRAIL-R1/TNFSF10A) and DR5 (TRAIL-R2/TNFSF10B) in humans, and TRAIL-R (MK/mDR5) in mice. TRAIL is found in Leydig cells and germ cells during development and TRAIL-R is predominantly expressed in post-meiotic germ cells in the rat. The major role of TRAIL in male reproduction is still unclear. In this study, we investigated TRAIL-/- mice and evaluated the possible role of TRAIL in germ cell development by measuring testis weight, germ cell apoptosis, and spermatid head count in periuberal (28 day-old) or young adult animals (44 day-old). Our results revealed that there was a significant difference in testis to body weight ratio between C57 and TRAIL-/- mice. Also, periuberal TRAIL-/- mice show a dramatic increase in the basal germ cell apoptotic index (A.L. 21.54%) as compared to the C57BL6/J wild-type strain (5.16%). The A.I. in adult C57 mice had dropped to 1.5%, but remained elevated in adult TRAIL-/- mice (20.2%); indicating a sustained high incidence of germ cell apoptosis. Spermatid head counts in adult TRAIL-/- mice were dramatically reduced compared with C57 (39%), indicating these animals suffer a marked decline in the production of mature spermatzoa. We hypothesize that TRAIL is an important factor for maintaining germ cell homeostasis during germ cell development via a death receptor-dependent mechanism.

1713 Investigation of Sex-Dependent Toxicity of Cysteinyi Leukotriene Receptor 1 Antagonist Zafirlukast.


The FDA-approved prescription drug label states zafirlukast, a cysteinyl leukotriene receptor 1 (CysLT1) antagonist, induces adverse hepatic events "predominantly in females." To model this possible sex-dependent toxicity, we examined suspension culture rat hepatocytes from female rats were more sensitive than male rats in all three parameters. Changes in ATP and cellular respiration were seen as early as 1 hr while cytochrome oxidase activity was not affected. Hepatocytes from male rats were more sensitive than male rats in all three parameters. Changes in ATP and cellular respiration were seen as early as 1 hr while cytochrome oxidase activity was not affected. In human lung BEAS-2B cells, both compounds induced reactive oxygen species (ROS) production, while caspase 3/7 activity was not affected. The compounds caused little/less H2O2, and oxidative DNA damage. Hence, a modulation of this process may significantly influence the cytoxicity and pro-death effects of the compounds.

1714 1,3-Dinitropropene (1,3-DNP)- and 1,8-DNP-Induced DNA Damage and Cell Death: Possible Link to Carcinogenic Effects.

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Nitro-polyacrylic aromatic hydrocarbons (nitro-PAHs) are found on particulate matter from diesel and gasoline exhaust. Often they are found to have greater mutagenic and carcinogenic potentials when compared with their parent PAHs. In the present study, we have compared the genotoxic and cytotoxic effects of two closely related carcinogenic dinitropropenes (DNPs), 1,3-DNP and the more potent 1,8-DNP. In human lung BEAS-2B cells, both compounds induced reactive oxygen species (H2O2; flow cytometry), oxidative DNA damage (comet assay), and DNA damage response measured as phosphorylation of p53 (Western blotting) at non-cytoxic concentrations (3-30 μM). In mouse hepatoma Hepa1c1c7 cells, 1,3-DNP (>3 μM) induced cell death (a mixture of apoptosis and necrosis), while 1,8-DNP caused necrotic effects. The compounds caused little/less H2O2, and oxidative DNA damage that in BEAS-2B. Interestingly, 1,8-DNP was overall more potent than 1,3-DNP with regard to the induction of single-strand DNA breaks (comet assay) and the formation of DNA adducts (32P-postlabelling). Furthermore, 1,8-DNP gave a stronger DNA damage response (phosphorylation of H2AX and p53) than 1,3-DNP in Hepa1c1c7 cells. Thus, there was no apparent link between the induction of cell death and early increases in ROS-formation or DNA damage/DNA damage responses in the two cell lines. 1,3-DNP-induced apoptosis in Hepa1c1c7 cells was specifically associated with mitochondrial damage (increase formation of superoxide anion; flow cytometry), but was also dependent on the p53-linked transcriptional apoptotic pathway (inhibitors and siRNA). We suggest that the stronger carcinogenic potency of 1,8-DNP compared to 1,3-DNP is due to its greater DNA damage properties, which in combination with its lower potency to induce cell death increases the probability of causing mutations.

1715 Role of NOX Mediated Autophagy in Reducing Cytotoxic Effects of Erlotinib in Head and Neck Cancer Cells.

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Most head and neck squamous cell carcinomas (HNSCC) overexpress Epidermal Growth Factor Receptor (EGFR) which makes it an attractive candidate for molecular targeted therapies. A combination of surgery, radiation and chemotherapeutic agents like EGFR inhibitors are routinely used in the treatment, however, many HNSCC tumors become resistant to EGFR inhibitors. The cellular self-degradation process autophagy is activated by oxidative stress and has recently been reported to reduce the efficacy of chemotherapy. Previous work in our lab has shown that the EGFR inhibitor Erlotinib induces oxidative stress via NADPH Oxidase 4 (NOX4) in HNSCC cells. The purpose of this study is to determine if Erlotinib induces autophagy in HNSCC cells via NOX4 and if autophagy is a pro-survival or pro-death mechanism. Erlotinib induced cytotoxicity (as determined by clonogenic assay) in FaDu and Cal-27 HNSCC cells compared to control treated cells. Erlotinib stimulated the expression of the autophagy marker LC3B-II in both cell lines as determined by western blot and immunofluorescence assays. Knockdown of autophagy genes Beclin-1 and Atg5 sensitized both cell lines to the cytotoxic effect of Erlotinib, suggesting that autophagy may be a pro-survival mechanism. Erlotinib increased NOX4 mRNA and protein levels in FaDu and Cal-27 cells. Treatment with DPI (diphenyleneiodonium) and a p58 inhibitor in the presence of Erlotinib suppressed the increase in LC3B-II expression in FaDu and Cal-27 cells. Finally, knockdown of NOX4 using adenoviral siNOX4, partially suppressed the activation of LC3B-II in FaDu and Cal-27 cells. These results suggest that Erlotinib may activate autophagy in HNSCC cells as a pro-survival mechanism, and NOX4 may play a role in mediating this effect. In conclusion, NOX4-induced autophagy may play a role in reducing the efficacy of Erlotinib. [Supported by NIH grant K01-CA134941 and ACS grant IRG-77-004-34].

1716 The Aryl Hydrocarbon Receptor (AhR) Suppresses Apoptosis in UVB-Irradiated Keratinocytes and May Serve As a New Target for Chemoprevention.

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Exposure of keratinocytes to ultraviolet (UV) radiation results in the initiation of apoptosis, a protective mechanism that eliminates cells harbouring irreparable DNA damage. Hence, a modulation of this process may significantly influence the initiation and progression of UV-induced skin cancer. We have found that the aryl hydrocarbon receptor (AhR), a ligand-activated and UV-sensitive transcription factor, serves an anti-apoptotic function in UVB-irradiated human keratinocytes. Chemical and siRNA-mediated disturbance of AhR signaling, significantly enhanced UVB-induced apoptosis. This effect was due to a loss of expression of EEF2 and its downstream target checkpoint kinase-I (CHK1), two factors critical for cell-cycle control and DNA damage response. Ectopic overexpression of EEF2 in AHR-knockdown keratinocytes restored CHK1 expression and diminished the observed sensitization to UVB-induced apoptosis. Accordingly, experimental CHK1 recovery alone was also sufficient to attenuate UVB-induced apoptosis in AHR-knockdown keratinocytes, indicating that that the loss of proper checkpoint control drives damaged keratinocytes into programmed cell death. Our results demonstrate for the first time an interplay between AHR, EEF2 and CHK1 and identify this signaling axis as a novel anti-apoptotic pathway in keratinocytes, which may represent a putative target for chemoprevention of non-melanoma skin cancer.

1717 Role of Secretory Phospholipase A2 in the Toxicity of Bile Aides to Prostate Cancer Cells.

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Bile acids mediate the digestion and absorption of fats and fat-soluble vitamins; however, pathological increases are associated with choleostasis and cell death. Recent studies show that high concentrations of bile acids can induce apoptosis in...
several cells, including cancer cells, by mechanisms that are not fully understood. The goal of this study was to determine the toxicity of three different bile acids (chenodeoxycholic acid, deoxycholic acid, and lithocholic acid) in prostate cancer cell lines (PC-3, LNCaP; and DU-145). Treatment of cells with bile acids induced time- and concentration-dependent decreases in MTT staining, a marker of cytotoxicity, with IC50 values of approximately 100-200 μM after 72 hr. In general, lithocholic acid was more potent than chenodeoxycholic acid, followed by deoxycholic acid. Further, LNCaP cells tended to be more susceptible to bile acid-induced toxicity, than either DU-145 or PC-3 cells. Based on reports that bile acids increase the expression of inflammatory enzymes called secretory phospholipase A2 (sPLA2), we tested the hypothesis that these enzymes regulate the mechanisms of bile acid-induced cell death. Analysis of sPLA2 expression using quantitative PCR showed that several sPLA2 isoforms were expressed in PC-3, LNCaP and DU-145 cells, including Group IB, II A, V and X sPLA2. Nevertheless, treatment of cells with the sPLA2 inhibitor LY317275, prior to exposure to bile acids, did not alter MTT staining compared to cells exposed to bile acids alone. Similar results were seen with the calcium-independent PLA2, (IPLA2) inhibitor bromelain lactone. Collectively, these data show the novel finding that bile acids can induce toxicity to prostate cancer cells and suggest the inhibition of sPLA2, nor IPLA2, activity mediate the mechanisms of cytotoxicity.

1718 Etoposide-Induced Mitochondria-Dependent Apoptosis through C-Jun N-Terminal Kinases, Extracellular Signal-Regulated Kinases, and Glycogen Synthesis Kinase-3β/α Pathway in Pancreatic β-Cells.

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Etoposide, a semisynthetic derivative of podophyllotoxin, is an important chemotherapeutic agent and widely used to treat human cancers. Etoposide also can produce the severe side effects and cause cell damages and the physiological dysfunctions. However, the toxicological effects of etoposide-induced pancreatic β-cell death remain unclear. Here, we investigate the cytotoxic effect and its possible mechanisms of etoposide on pancreatic β-cells. Treatment of pancreatic β-cell-derived RIN-m5F cells with etoposide (1-100 μM) for 24 h significantly reduced cell viability and underwent apoptosis, accompanied with mitochondrial dysfunctions. Moreover, etoposide triggered the protein phosphorylation of glycogen synthesis kinase (GSK)-3(β)/α at 8 h treatment and maintained to 24 h, which could be reversed by lithium chloride (LiCl, a specific inhibitor of GSK-3(β)/α). In addition, etoposide (20 μM) markedly increased the phosphorylation of JNK and ERK1/2, but not p38. Pharmacological inhibitors SP600125 and PD98059 effectively attenuated etoposide-induced caspase-3 activity and JNK and ERK1/2 activation, but LiCl could not reverse the phosphorylation of JNK and ERK1/2 induced by etoposide. In conclusion, this results suggest that etoposide exerts its cytotoxicity on pancreatic β-cells by inducing the mitochondria-dependent apoptosis through JNK/ERK activation-regulated GSK-3(β)/α signaling pathway.

1719 Berberine Induces Human Tongue Squamous Carcinoma Cell Apoptosis through PI3K-Regulated ER Stress Pathway.

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Until now, oral cavity squamous cell carcinoma (OSCC) is the most common head-and-neck cancer, which accounts for approximately 3% of all newly diagnosed cancer cases. Despite of recent advances in surgical, radiotherapy, and chemotherapy treatment protocols, it has been discussed that OSCC could not be eradicated. Berberine is a natural alkaloid. Recently, berberine has been showed to inhibit metastasis in lung cancer cells, and cytotoxicity in glioma, prostate and nasopharyngeal cancer cells. However, the therapeutic effects and the possible mechanisms of berberine in OSCC are still unclear. Results found that berberine significantly decreased cell viability in human tongue squamous carcinoma derived SAS cells line. Besides, berberine increased the ER-stress signals, including Grp94, CHOP Xbp-1 mRNAs and proteins. Further, the pro-caspase-12 protein level was decreased, while the caspase-12 mRNA level was increased. In the mitochondrial pathway, berberine increased Bax, Bak, Bid mRNAs and proteins levels and decreased Bcl-2 mRNA and proteins levels. Besides, the pro-caspase-9, pro-caspase-7, and pro-caspase-3 protein levels was decreased. We also found that phospho-ATP protein level was decreased after berberine treatment in cells. Besides, L294002, the inhibitor of phosphoinositide 3-kinases (PI3Ks), promoted ER-stress and mitochondrial related apoptosis signals. Through this study, suggested that berberine may be a useful compound in inhibition of OSCC through PI3K-ATK regulated ER-stress and mitochondrial pathways.

1720 Amiodarone-Induced Autophagy Protects Lung Epithelial Cancer Cells from Death.

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Amiodarone-induced toxicities in various off-target organs including cornea, liver, skin, thyroid and lung often limit the use of this drug especially in the case of long-term treatment. Despite low incidence rate, amiodarone-induced pulmonary toxicity is the one of greatest concern and leading cause for discontinuation. In the present study, we investigated the induction of autophagy following amiodarone treatment in vitro and the relevance of autophagy to the pulmonary toxicity. Amiodarone treatment caused morphological change in H460 cells. Increased numbers of cytoplasmic vacuoles as well as round and floating cells were observed. Amiodarone-induced autophagy was demonstrated by increased LC3-II to LC3-I conversion, upregulation of Atg5 and Atg7 and dephosphorylation of p62 protein. Autophagic flux as determined by lysosomal inhibitor bafilomycin A1 was also increased by amiodarone. The phosphorylation of AMP-activated protein kinase at Thr172 was increased and that of Ake at Ser73 was decreased by amiodarone which resulted in the reduction of mTOR and p70S6 kinase phosphorylation. To determine the role of autophagy in amiodarone toxicity, amiodarone-induced cell death was evaluated in the presence of 3-methyladenine or by knocking down autophagy related genes, Atg5 or Atg7. Inhibition of autophagy decreased the cell viability and increased apoptosis significantly. In conclusion, amiodarone-induced autophagy functions to rescue stressed cells from death. The findings have important implications in developing further strategy to minimize drug-induced toxicity in human.

1721 Paraquat Causes Hepatocytes Death via Oxidative Stress-Induced JNK/ERK Activation Regulated Mitochondria-Dependent Apoptosis Pathway.

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Paraquat (1,1'-dichloro-2,2'-bipyridyl dichloride, PQ), a common herbicide used all over the world, is toxic to human beings and causes severe injuries to multiple organs, including lung and liver. However, the toxicological effects and molecular mechanisms of PQ-induced on hepatocytes are mostly unclear. In this study, we found that PQ significantly reduced the cell viability in rat hepatocytic cell line H4-II-E cells. Treatment of H4-II-E cells with PQ also induced several features of mitochondria-dependent apoptotic signals, including loss of mitochondrial membrane potential (MMP), increase in cytosolic cytochrome c release, activation of PARP and caspase-3/7, and increased oxidative stress injuries such as reactive oxygen species (ROS) generation and glutathione depletion. These PQ-induced apoptotic-related signals could be effectively reversed by antioxidant NAC. Moreover, PQ increased the phosphorylation of JNK and ERK1/2, but not p38. Pharmacological inhibitors SP600125, PD98059, and NAC significantly attenuated PQ-induced cytotoxicity, caspase-3/7 activation, MMP loss, and inhibited death.
the phosphorylation of JNK and ERK1/2. Taken together, these results suggest that PQ exerts its cytotoxicity on hepatocytes by inducing apoptosis via an oxidative stress-induced JNK and ERK1/2 activation regulated mitochondria-dependent signaling pathway.

1722 Oxidative DNA Damage and Apoptosis Induction after Enniatin B Exposure in Caco2 Cells.


Enniatin (EN) B is a cyclohexadepsipeptidic mycotoxin produced by Fusarium spp. often found in cereals and cereal-based products. Its cytotoxic potential has been reported but the mechanisms involved in its toxicity remain to be elucidated. Since the oxidative pathway could be implicated in mycotoxin's toxicity, in this study the generation of reactive oxygen species (ROS) and lipid peroxidation (LPO) have been investigated in human colorectal adenocarcinoma (Caco-2) cells. Subsequently the induction of oxidative stress have been assumed to be directly related to DNA damage and apoptosis. Cells were exposed to EN B at sub-cytotoxic concentration of 1.5 and 3 μM. A significant increase (p<0.05) in ROS production was observed by the fluorescent probe H2-DCFDA after 3 μM exposure in Caco-2 cells from 10 up to 120 min. LPO was determined by thiobarbituric acid reactive substances (TBARS) after 24 h of exposure. Significant increase of malondialdehyde (MDA) was observed after the highest (3 μM) concentration tested of 48% (p<0.05), as compared to control. Genotoxicity was evaluated through the alkaline Comet assay after 2 and 24 h of exposure. Median tail moment (μM) was significantly high respect to the control with a dose-dependent relationship (p<0.05) after short and longer exposure. The induction of apoptosis and necrosis was assessed by flow cytometry. Annexin V coupled to FITC in combination with PI was used to determine different apoptotic phases after 24 h of exposure to EN B (1.5 and 3 μM). Both concentrations induced apoptosis and necrosis in a dose-dependent manner. These findings show that 3 μM concentration of EN B caused oxidative damage, by means of ROS generation that could be responsible of LPO. DNA damage and triggers apoptosis in Caco2 cells.

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1723 Mevalonate Pathway Plays a Major Role in Adriamycin Resistance.

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In a search for novel mechanisms of resistance to adriamycin, an anthracycline anti-tumor antibiotic used in cancer chemotherapy, we have previously screened a ORF library derived from budding yeast for genes related to adriamycin resistance and found that overexpression of the gene for HMG-CoA synthase, an enzyme in mevalonate pathway, confers resistance to adriamycin in budding yeast. We have also found that promotion of mevalonate pathway decreased the toxicity of adriamycin in yeast cells.

In this study, we examined the relationships between enzymes involved in mevalonate pathway and the adriamycin resistance. And then that deletion of the gene for Ram1, a farnesyltransferase, reduced the degree of adriamycin resistance induced by overexpression of HMG-CoA synthase. These results suggest that overexpression of HMG-CoA synthase could decrease the adriamycin toxicity through promotion of farnesylation of proteins. Moreover, overexpression of HMG-CoA synthase or addition of mevalonate to culture medium decreased the toxicity of adriamycin in human breast MCF7 cells, suggesting that mevalonate pathway plays a key role in mechanism of adriamycin resistance not only in yeast cells but also in human cells.

1724 Carbamazepine Suppresses Ischemia/Reperfusion Injury to Mouse Livers by Enhancing Autophagic Flux.


BACKGROUND: Onset of the mitochondrial permeability transition (MPT) plays a causative role in ischemia/reperfusion (I/R) injury, a pathological event occurring during organ transplantation, cardiac failure and hemorrhagic shock. Current therapeutic strategies for reducing reperfusion injury remain disappointing. As autophagy selectively and timely eliminates abnormal or damaged cellular constituents and organelles such as dysfunctional mitochondria, this lysosome-mediated catabolic process confers cytoprotection against I/R injury and various diseases. We have shown that calpain-dependent depletion of autophagy-related proteins (Atg) causes the MPT and hepatocyte death after I/R. Carbamazepine (CBZ), an FDA approved anticonvulsant drug, has recently been reported to increase autophagy. The AIM of this study was to investigate the effects of CBZ on hepatic I/R injury.

METHODS: Hepatocytes and livers from male C57BL/6 mice were subjected to simulated and in vivo I/R, respectively. Cell death, intracellular calcium, calpain activity, changes in Atg, autophagic flux, MPT and mitochondrial membrane potential after I/R were analyzed in the presence and absence of 20 μM CBZ.

RESULTS: CBZ significantly increased hepatocyte viability after reperfusion, as judged by propidium iodide fluorometry. Confocal microscopy of rhod-2, fluo-4, calcine and tetramethylrhodamine methylester revealed that CBZ prevented reperfusion-induced mitochondrial calcium loading, onset of the MPT and mitochondrial depolarization. Immunoblotting and fluorometric analysis showed that CBZ blocked calpain activation, Atg6 depletion, and loss of autophagic flux after reperfusion. Intravital multiphoton imaging of anesthetized mice demonstrated that CBZ substantially reversed autophagic defects and mitochondrial dysfunction after I/R in vivo.

CONCLUSION: CBZ protects hepatocytes against I/R injury by preventing a temporal sequence of calcium overloading, calpain activation, Atg6 depletion, defective autophagy, onset of the MPT, and cell death.
Characterization of Cigarette Smoke and Menthol on Human Alveolar Adenocarcinomic (A549) Cells.

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Purpose: Menthol provides local anesthesia to nerve endings, allowing smokers to take deeper, longer inhalations, thus giving longer exposure to carcinogenic elements found in cigarette smoke. There are few cellular studies addressing evidence concerning the risk of developing lung cancer between mentholated and non-mentholated cigarettes. We would like to determine if an association exists between mentholated cigarettes and incidence of lung cancer in the smoker population.

Methods: Using an in vitro model, we aimed to characterize the effects of cigarette smoke, with or without menthol, in human alveolar adenocarcinomic (A549) cells and determine if the rate of cell proliferation and cell death is altered.

Results: Data suggest that menthol alone decreased cell viability and increased Annexin positive cells while co-treatment with CSC + menthol did not affect cell viability. Pan-caspase activation was not significantly altered for CSC, menthol, or CSC + menthol. Interestingly, cells treated with CSC only appeared to emit a source of auto fluorescence. Future studies are required to address the issue.

Apoptosis is a programmed form of cell death executed by caspases (cysteine proteases) that cleave substrates exclusively after aspartic acid residues. In response to stress, cells often activate the intrinsic apoptotic pathway, wherein mitochondrial (mitochondrial proteins, such as cytochrome c and Smac, into the cytosol. In particular, cytochrome c promotes formation of a caspase-activating complex known as the Apaf-1-caspase-9 apoptosisosome. Heat shock (HS) is an ancient stress that activates both prosurvival (thermotolerance) and prodeath cellular responses, and we have previously shown that HS induces apoptosis through pathways that involve MOMP and downstream effector caspase-3 activation, but do not require activation of the apoptosisosome. HS-induced MOMPs are strictly controlled by BCL-2 family members, which include both pro-apoptotic (e.g., Bax, Bid, Bcl, etc.) and anti-apoptotic (e.g., Bcl-2, Mcl-1, etc.) proteins. Other studies suggest that Bim induces MOMP following cleavage of BID by caspase-2. However, we observe that mouse embryonic fibroblasts (MEFs, lacking either caspase-2 or BID, remain sensitive to HS. In contrast, we find that bim-/- MEFs are highly resistant to HS-induced apoptosis and exhibit significantly decreased levels of MOMP. Thus, our findings indicate that Bim is essential for HS-induced death, while caspase-2 and BID function as part of an amplification loop. Interestingly, Bim is known to induce MOMP through direct activation of Bax and/or Bak, but we find that bax-/-/bak-/- MEFs are only partially resistant to HS-induced cell death, implying the existence of a Bim-independent, Bax/Bak-independent pathway. We speculate that another proapoptotic Bcl-2 family, namely Bok, may functionally substitute for Bax and Bak, or alternatively that Bim may directly permeabilize lysosomal membranes, resulting in the release of proapoptotic cathepsins. (These studies were supported by grants, CA129521 and GM096101, from the NIH.)
apoptosis whereas IL-6 siRNA abolished the protective effect of SDF-1β. CXCR7 siRNA, but not CXCR4 antagonist abolished SDF-1β’s protective effect and above related signal transduction. These in vitro results suggest that SDF-1β prevents palmitate-induced cardiac apoptosis via its receptor CXCR7 and further activating AMPK-mediated IL-6 excetration. In vivo studies, by using type 2 diabetes models, we confirmed that high fat diet induced cardiac apoptosis, and that SDF-1β prevented high fat diet-induced cardiac apoptosis along with its activation of AMPK. This important finding opens a new road for the research of SDF-1β’s cardiac protection that is irrelevant with its well-known function of stem cell mobilization.

**1732** Paraxon Induces Cell Death by the Activation of Caspases in Cultured Human Pulmonary Cells.

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INTRODUCTION: Organophosphate chemicals are known to induce pulmonary toxicity in both humans and experimental animals. To elucidate the mechanism of organophosphate-induced cytotoxicity, we examined the effects of parathion and paraoxon on primary cultured normal human bronchial epithelial cells (NHBECs) as well as small airway cells (SAECs).

METHODS: We evaluated the viability/cytotoxicity of primary NHBECs and SAECs following exposures to the organophosphate chemicals parathion and paraoxon. Since caspases have a major role in the regulation of apoptosis and cell death, we evaluated paraoxon-induced cell death in the presence of the caspase inhibitor Z-VAD.

RESULTS: Paraoxon failed to induce losses in cellular viability at concentrations known to cause toxicity in vivo following a 24 hour treatment (IC50 >9mM) in either NHBECs or SAECs. Its toxic metabolite, paraoxon, produced a dose-dependent decrease in cell viability following a 24 hour treatment period for both pulmonary cell types (IC50 ~0.5mM). Treatment with parathion also induced caspase activation in these cells. Pharmacological inhibition of caspases with Z-VAD during parathion treatment protected against paraoxon-induced cell death.

CONCLUSIONS: These combined data suggest that paraoxon induces cell death in NHBECs and SAECs, at least in part, through the activation of caspases.

**1733** BDE-154 Decrease Cell Proliferation, Viability and Mitochondrial Membrane Potential Leading to HepG2 Cell Death.

A. O. Souza1, L. C. Pereira1, M. Tasso1, D. P. Oliveira2 and D. I. Dotta1.

INTRODUCTION: Polybrominated diphenyl ethers (PBDEs) are a class of flame retardants composed by 209 different congeners with evidences of several toxic potential on the organisms. Toxic damages of lower and higher brominated congeners have been widely reported but more information about other congeners to clarify risk of PBDEs class on the health is still necessary. OBJECTIVE: The aim of this work was to investigate the effects of the BDE-154 on HepG2, because it had showed growth levels of bioaccumulation on the biotic compartments. METHODOLOGY: We performed a cell proliferation test using SRB colorimetric wash liquid had the same electrical conductivity as pure H2O, indicating very few wash liquid had the same electrical conductivity as pure H2O, indicating very few.

RESULTS: Our results showed that BDE-154 are capable of inhibiting cell proliferation at doses starting at 10μM (p<0.05) after 48 hours of exposure, although the results of MTT assay had shown a significant decrease in cell viability on lower doses in both times of incubation. Changes on cell proliferation and viability were followed by decrease of MPP and an increase of phosphatidylserine exposure at the cell membrane. CONCLUSIONS: According to our results, BDE-154 presents a significant potential to interfere with the mitochondrial homeostasis of HepG2 cells which can lead to apoptosis in the concentration of 25 μM.

**1734** Computational Dosimetry Driven Hazard Ranking of 25 Metal Oxide Nanomaterials Using Low- and High-Throughput In Vitro Toxicity Data.

D. Telecsa1, J. G. Teegarden1, T. Xia4, H. Zhang1, A. Nel3, J. G. Pounds1 and B. D. Thrall1,2.

RESULTS: Our results shown that BDE-154 are capable of inhibiting cell proliferation, and that SDF-1β prevented high fat diet-induced cardiac apoptosis along with its activation of AMPK.

**1735** Ranking and Profiling Nanomaterial (NM) Bioactivity by ToxCast High-Throughput Screening (HTS).


INTRODUCTION: Organophosphate chemicals are known to induce pulmonary toxicity in both humans and experimental animals. To elucidate the mechanism of organophosphate-induced cytotoxicity, we examined the effects of parathion and paraoxon on primary cultured normal human bronchial epithelial cells (NHBECs) as well as small airway cells (SAECs).

METHODS: We evaluated the viability/cytotoxicity of primary NHBECs and SAECs following exposures to the organophosphate chemicals parathion and paraoxon. Since caspases have a major role in the regulation of apoptosis and cell death, we evaluated paraoxon-induced cell death in the presence of the caspase inhibitor Z-VAD.

RESULTS: Paraoxon failed to induce losses in cellular viability at concentrations known to cause toxicity in vivo following a 24 hour treatment (IC50 >9mM) in either NHBECs or SAECs. Its toxic metabolite, paraoxon, produced a dose-dependent decrease in cell viability following a 24 hour treatment period for both pulmonary cell types (IC50 ~0.5mM). Treatment with parathion also induced caspase activation in these cells. Pharmacological inhibition of caspases with Z-VAD during parathion treatment protected against paraoxon-induced cell death.

CONCLUSIONS: These combined data suggest that paraoxon induces cell death in NHBECs and SAECs, at least in part, through the activation of caspases.
Coating Nanoporous Alumina: Cellular Responses to Surface Layers of Zinc and Titanium Oxides.

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Nanoporous anodic aluminum oxide (AAO) membranes have applications in skin wound repair; the surface topography of these materials has been shown to modulate wound healing. Previous work by our group has shown that coating titanium dioxide (TiO2) onto nanoporous AAO does not affect cell viability and that zinc oxide (ZnO)-coated AAO membranes exhibit antimicrobial activity against several bacterial strains associated with skin infection. In this study, the nanoporous structure of 20, 100, or 200 nm pore diameter AAO membrane substrates was maintained by depositing 5-10 nm-thick layers of TiO2 or ZnO using an atomic layer deposition (ALD) process involving alternative adsorption/hydrosilylation of Ti isopropoxide or diethyl Zn, respectively. X-ray spectroscopy (EDX) confirmed the presence of ZnO or TiO2 on AAO membranes. Cell viability and inflammatory responses were evaluated on ZnO- and TiO2-coated AAO membranes. RAW 264.7 macrophages cultured on ZnO-coated AAO exhibited a significant decrease in metabolic activity (MTT reduction) after 24 or 48 h exposure; however, no changes were observed in lactate dehydrogenase (LDH) release or reactive oxygen species (ROS) production. L929 fibroblasts grown on TiO2-coated AAO had higher viability (MTT assay) than controls (uncoated AAO) after 48 h; no change in viability was observed using the Neutral Red assay. No significant increase in fibroblast cell proliferation (DNA assay) was observed. No TNF-α production or decrease in viability was seen in RAW 264.7 macrophages grown on TiO2-coated membranes. The total adsorbed protein per unit area on uncoated and TiO2-coated AAO was up to 12-times greater than that on tissue culture polystyrene. The results show that nm-thick ALD coatings maintain the nanoporous structure of AAO membranes. Further, antimicrobial ZnO coatings decrease cell viability and TiO2 coatings appear to be more biocompatible in vitro.

Potential Phototoxicity of Al(OH)3-Coated TiO2 Nanoparticles in Retinal Pigment Epithelial Cells.


Titanium dioxide (TiO2) nanoparticles (NPs) exposed to UV radiation generate reactive oxygen species (ROS). As a component of sunscreen formulations, TiO2 NPs may be coated with Al(OH)3 to prevent ROS from causing oxidative damage to tissues. Simulated swimming pool water (SSPW) degrades the Al(OH)3 coating which could reduce the coating’s protective function. We examined the potential phototoxicity of 25 nm, Al(OH)3-TiO2 NPs that had been aged in SSPW for 0, 5, 50, and 175 days. The results show that nano-phototoxicity of TiO2 Anatase Nanoparticles in B6C3F1 Male Mouse Pig-A and Flow Cytometric Micronucleus Assays.

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In vivo micronucleus and Pig-a (phosphatidylidyinositol glycan, class A gene) mutation assays were conducted to evaluate the genotoxicity of 10 nm titanium dioxide anatase nanoparticles (TiO2-NPs) in mice. Groups of five 6- to 7-week-old male B6C3F1 mice were treated intravenously for three consecutive days with 0.5, 5.0, and 50 mg/kg TiO2-NPs for the two assays; mouse blood was sampled one day before the treatment and on Day 4, and Weeks 1, 2, 4, and 6 after the beginning of the treatment; Pig-a mutant frequencies were determined at Day-1 and Weeks 1, 2, 4 and 6, while percent micronucleated-erytocyte (%MN-RET) frequencies were measured on Day 4 only. Additional animals were treated intravenously with three
daily doses of 50 mg/kg TiO2-NPs for the measurement of titanium levels in bone marrow after 6, 24, and 48 hrs of the last treatment. The measurement indicated that the accumulation of the nanoparticles reached the peak in the tissue 4 hrs after administration and the levels were maintained for a few days. No increase in either Pig-a or mouse reticulocyte was detected, although the RETs were reduced in the treated animals on Day 4 in a dose-dependent manner indicating cytotoxicity of TiO2-NPs in the bone marrow. These results suggest that although TiO2-NPs can reach the mouse bone marrow and are capable of inducing cytotoxicity, the nanoparticles are not genotoxic when assessed with in vitro micronucleus and Pig-a gene mutation tests.

**1744 Inhibition of Adipogenic Differentiation of Human Mesenchymal Stem Cells by TiO2 Nanoparticles.**

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Nanotechnology has resulted in the creation of many new materials and devices with a vast range of applications. TiO2 has historically been used as a pigment with many applications, and nanoscale TiO2 is additionally appearing in human consumer products, raising safety issues of human health and environmental concerns. Stem cells are proposed to be attractive tools for toxicity testing because of their sensitivity to external stimuli during differentiation. The physical and chemical properties of TiO2 nanoparticles were characterized using transmission electron microscopy (TEM), nanoparticle tracking analysis, dynamic light scattering, Brunauer-Emmett-Teller, Raman and X-ray fluorescence spectroscopy. We examined the impact of TiO2, and other nanomaterials, on cytotoxicity and differentiation of human mesenchymal stem cells (hMSC). TiO2 nanoparticles induced cytotoxicity in a concentration-dependent manner in hMSCs. Additionally, the differentiation of hMSCs to adipocytes, determined using imaging and Oil Red O staining, was inhibited in a concentration-dependent manner. Moreover, uptake of the TiO2 nanoparticles by hMSCs was confirmed using TEM, suggesting a possible endocytosis pathway. The mRNA expression for the adipogenic markers adiponectin, aP2, and LPL were significantly reduced to 61%, 53%, and 61% of control levels following exposure to TiO2 nanoparticles. Furthermore, incubation of the TiO2 with the mesenchymal stem cell line resulted in the identification of 208 proteins associated with the nanoparticles using proteomic and mass spectrometry analyses. These results indicate (i) the interaction and impact of nanoparticles with stem cells is selective (other nanoparticles did not induce this effect), (ii) TiO2 nanoparticles inhibited differentiation of mesenchymal stem cells, and (iii) further work is needed to elucidate the mechanism of action of TiO2 in this cell population.
Mechanisms of Silica Nanoparticle-Induced Interleukin-8: Requirement of p38/TACE/TGF-α/EGFR-Cascade and NF-κB Signalling in Lung Epithelial Cells.

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Nanoparticles (NPs) of non-crystalline (amorphous) silica particles (SiNPs) are used in a large range of products. Inhalation of NPs represents a potential health hazard and may induce inflammation in lung tissues. We have previously shown that SiNPs induced marked cytokine responses independently of particle uptake in human bronchial epithelial cells (BEAS-2B). In the present study the mechanisms involved in SiNP-induced IL-8 responses were further examined. SiNP-exposure induced an early increase in phosphorylation of p65 (NF-κB) as well as the three main MAP-kinases ERK1/2, p38 and JNK, concurrent with an early up-regulation of IL-8 mRNA. SiNPs also induced a time-dependent increase in phosphorylation of the epidermal growth factor receptor (EGFR) and release of the EGFR-ligand transforming growth factor (TGF)-α. SiNP-induced IL-8 responses were attenuated by the p38-inhibitor SB202190, the NFκB-inhibitor PDT-p65 and siRNA against p65, as well as the EGFR-inhibitor AG1478, a TGF-α-neutralizing antibody and TAPI-1 (inhibitor of the metalloprotease TACE which cleaves pro-TGF-α to TGF-α). However, inhibitors of ERK and JNK did not exert any effect on SiNP-induced IL-8. Moreover, SiNP-induced EGFR-phosphorylation was inhibited by AG1478 and TAPI-1, and SB202190 reduced the SiNP-induced TGF-α response. The SiNP-induced phosphorylations of p38 and p65 were not affected by TAPI-1 or AG1478. Thus, SiNP appeared to induce EGFR-phosphorylation through a p38- and TACE-dependent cleavage/release of TGF-α. Interestingly, EGF and TGF-α induced little effect on IL-8 release compared to SiNP suggesting that EGFR-signalling alone is an insufficient stimuli for IL-8 induction. In conclusion, SiNP-induced IL-8 responses seemed to require activation of p38/TACE/TGF-α/EGFR-cascade, presumably acting in concert with the classical NF-κB pathway in BEAS-2B cells.

Role of Tungstate Nanoparticles in the Production of ROS and Induction of Cellular Damage.

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Alkaline-earth metal tungstate AWO3 (A = Ca, Ba, Sr) nanoparticles are currently being used in a variety of applications including use as components of medical equipment, optical fibers, gas sensors, and scintillator detectors. Due to tungstate nanoparticle versatility, their manufacturing is expected to increase within the next 10 years. Our ongoing study is designed to examine the effects of tungstate nanoparticle exposure in order to develop safe workplace practices to limit exposure. Electron Spin Resonance (ESR) was used to measure hydroxyl radical (OH) production of tungstate nanoparticles following incubation with either hydrogen peroxide (H2O2) or RAW 264.7 cells. Additionally, enhanced dark field microscopy was used to assess nanoparticle association and engulfment by RAW cells over multiple time points up to 3 hours. Assays measuring H2O2, production, oxygen consumption, DNA damage and lipid peroxidation were used to assess possible cellular injury following RAW cell incubation with tungstate nanoparticles. Data showed that tungstate nanoparticles are capable of producing OH in the presence of H2O2 and RAW cells. Further, tungstate nanowires produced significantly greater OH compared to nanospheres as shown through ESR measurements. Initial dark field microscopy data showed an increase in association between tungstate nanoparticles and cells and a decrease in non-specific binding after 3 hours of exposure. Cellular damage results in conjunction with ESR data will promote an understanding of tungstate nanoparticle toxicity. It is important to understand the damaging effects and free radical production produced by tungstate nanoparticle exposure in order to ensure that accurate toxicity models are developed and to promote proper training in nanoparticle inhalation prevention.
Long-Term Inhalation Study with Nanomaterials: Characterization of As-Produced Properties and Persistence Simulation Assays.

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The most urgent knowledge gap in nanosafety research concerns the long-term fate and effects of nanomaterials that may have been released into the air or the environment. BASF and the German Ministry for the Environment (BMU) have initiated a chronic inhalation study as part of the EU project NanoREG. Here we report whether the materials chosen can be regarded as representatives of poorly soluble biopersistent (PSB) particles.

The original batch of CeO2 distributed as NM212 from the OECD sponsorship program and an identical reproduction of the BaSO4 distributed as NM220 were characterized by 20 physical-chemical endpoints according to the nanospecific REACH guidance R7.1, benchmarked against TiO2 (NM105) and the non-persistent SiO2 (Aerosil 150). The as-tested solubility was simulated by incubation for 28d in buffer (PBS) or phagolysosomal fluid (PSF), or 1d in stomach (0.1n HCl). Analysis included released ions by atom spectroscopy, agglomeration by analytical ultracentrifugation and laser diffraction, morphology by electron microscopy (TEM, SEM), and selected area electron diffraction (SAD). The re-characterization of CeO2 NM212 and BaSO4 NM220 confirms their conformity with earlier work on the OECD batches by PROSPECT and NanoCare projects. The solubility of CeO2 NM212 is as low as TiO2 NM105. The positive control SO2 dissolves as expected. The solubility of BaSO4 NM212 in all media, including PSF, is vanishing against the positive control, and noticeable only in 0.1n HCl. All materials, incl. TiO2, undergo morphological changes by Ostwald ripening and recrystallization in PSF and HCl, but retain their crystallinity. Dissolution is 0 to 0.1% per month; extrapolated on the chronic inhalation study, CeO2 NM212 and BaSO4 NM220 remain 99% persistent.
using microscopy, cytototoxic testing, x-ray microanalysis, particle counting, and inductively coupled plasma optical emission spectrometry. A nickel-chromium filament was coupled to a thicker silver coated copper wire. The silver coating was sometimes missing. Four tin solder joints attached the wires to each other and coupled the copper/silver wire to the air tube and mouthpiece. All cartomizers had evidence of use before packaging (burn spots on the fibers and electrophoretic movement of fluid in the fibers). Fibers in two cartomizers had green deposits that contained copper. Centrifugation of the fibers produced large pellets containing tin. Tin particles and tin whiskers were identified in cartridge fluid and outer fibers. Cartomizer fluid with tin particles was cytotoxic in assays using human pulmonary fibroblasts. The aerosol contained particles >1μm comprised of tin, silver, iron, nickel, aluminum, and silicate and nanoparticles (<100 nm) of tin, chromium, and nickel. Of 22 elements identified, 12 were present in concentrations higher than the minimum risk level. Many of the elements identified in EC aerosol are known to cause respiratory distress and disease. The presence of metal and silicate particles in cartomizer aerosol, often above minimal risk levels, demonstrates the need for improved quality control in EC design and manufacture and studies on how EC aerosol impacts the health of users and bystanders.

The interest in manufactured nanomaterials with potential new properties has led to increasing concern about their potential systemic uptake and fate as well as the associated risk to human health. Materials applied in processed food are of special interest. Synthetic amorphous silica (SAS) is a nanostructured material formed by flame hydrolysis or precipitation that has been used for decades. In commercial products, basic structural elements are submicron aggregates (fused nanosized primary particles) that themselves form micrometer (or even larger) agglomerates. SAS is employed in a variety of products, e.g. as free-flow agent in soup powders. As to risk evaluation, it is important to understand whether structural changes may occur during the further processing or after oral uptake. 

In a first step, we addressed possible effects of heating in water (modeled processing of soup powder) and acid environment (pH of gastric juice) on the structure and size distribution of SAS. Methods for the reproducible dispersion of SAS and the reliable determination of the volume weighted particle size distribution of SAS suspensions were developed and validated. Two food grade SAS types were studied: precipitated SAS and pyrogenic (fumed) SAS. SAS was first heated in water (100°C) and then poured into HCl to reach pH1-1.3 (paddle apparatus). During both steps time dependent changes in the volume weighted size distribution were monitored using laser diffraction (LD). 

LD with validation, e.g. by comparison to microscope analysis, proved as a reliable technique to characterize the dispersity of SAS suspensions and to evaluate the volume fraction of fine particles. Heating of SAS in water is only a weak dispersion leading to size distributions well above 1 μm. In acid environment (2 hours, pH=1.3) no significant changes in dispersity of SAS – neither agglomeration nor erosion of agglomerates or aggregates – was observed. In separate animal studies (rats, oral, repeated dose), no isolated nanosized primary particles of SAS were detected in the blood or organs.

Manufactured nanomaterials, as well as biomedicine. However, the correlation between exposure to metal nanoparticles and the increased incidence of cardiovascular disease remains elusive. The present study investigated the migration of monocytes and macrophage cholesterol uptake that are essential for atherosclerotic progression induced by nano-sized metal oxide particles. 

Zinc oxide nanoparticles have been widely used in industry, cosmetics, as well as biomedicine. However, the correlation between exposure to metal oxide nanoparticles and the increased incidence of cardiovascular disease remains elusive. The present study investigated the migration of monocytes and macrophage cholesterol uptake that are essential for atherosclerotic progression induced by nano-sized metal oxide particles. 

Methods and results: Human umbilical vein endothelial cells (HUVECs) were cultured and exposed to nano-sized TiO2 and ZnO. Exposure to ZnO for 6 hours reduced the cellular viability in a dose-dependent manner and increased the production of reactive oxygen species (ROS) whereas there were no changes by the exposure to TiO2. Exposure to ZnO for 21 hours increased the level of monocyte chemotactic protein-1 (MCP-1) in HUVECs, and cell migration of human monocytic leukaemia (THP-1) was observed after incubation with HUVEC supernatants. We also investigated the effect of nano-sized metal oxide particles on cholesterol uptake in THP-1 macrophages after stimulation with acetylated-LDL. The exposure to ZnO up-regulated the expression of membrane scavenger receptors of modified LDL particles and increased cholesterol uptake. 

Conclusion: The exposure to ZnO reduced the cellular viability in a dose-dependent manner and increased the production of ROS in HUVECs. The exposure to ZnO also induced THP-1 cell migration and increased macrophage cholesterol uptake. The study indicates that nano-sized ZnO particles accelerate foam cell formation in THP-1 macrophages.
Titannate nanomaterials (TiNMs) have been applied in various industrial products, including cosmetics, dye sensitized solar cells, and photocatalytic materials. There is also increasing concern on human health risk due to exposure to these nanomaterials during production and application of the products. Therefore, safety assessment of TiNMs is required. Several studies have been focused on toxicity and biological interactions of titannate nanoparticles (TiNPs), but less is known for titannate nanofibers (TiNFs). The aim of this study was to investigate the uptake patterns and toxicity of two TiNPs with different dimensions (about 50 and 70 nm in diameter and 500 and 1500 nm in length, respectively) compared with a spherical TiNP (about 170 nm in diameter) in human monococyte, THP-1 cells. The uptake of TiNMs into the cells was observed over the time for up to 24 hours using side scatter cytometry (SSC), which reflect internal cellular complexity, and electron microscopy. The results showed that all of TiNMs were immediately adsorbed on plasma membrane when they were exposed to the cells as observed by SEM and SSC. Envelopment of both TiNPs and TiNFs was observed by THP-1 cells occurred within 10 minutes exposure. Interestingly, SSC profile may suggest the different uptake patterns between TiNPs and TiNFs. In addition, the results from SSC and SEM suggested that TiNPs were taken up by the cells which took longer time than that of TiNPs because of their length. TiNPs with concentrations up to 100 μg/ml did not reduce cell viability, while TiNFs significantly reduced cell viability at concentration of 100 μg/ml. None of TiNPs (with concentrations up to 100 μg/ml) significantly trigger intra-cellular ROS generation. This information on cellular uptake and response might be useful for risk assessment of nanomaterials.

Metal-based nanoparticles (NPs) are generated from a variety of processes including welding, cutting and brazing. Airborne NPs are of particular concern over human exposure, as they can readily move in ambient air and enter the body through inhalation. The objective of this study was to investigate toxic effects on human alveolar type-II cells (A549) after air-delivery of various metal NPs. We used a spark discharge system (SDS) capable of generating and depositing airborne NPs directly onto cells at an air-liquid interface (ALI). The generated NPs (from various source materials including Cu, Zn electrodes and welding rods) were characterized by using a scanning mobility particle sizer, indcuently coupled plasma-mass spectroscope and electron microscopes (SEM, TEM). To better model in vitro repeated-low dose protocols we sequentially exposed lung cells to NPs in vitro (4 h exposure-2 h rest in an incubator-4 h exposure) and cell viability was determined by Alamar Blue assay at 4 h post-exposure. The SDS produced stable NP aerosols for 4 h (Cu, 8 x 10^6; Zn, 6 x 10^6; welding rods, 7 x 10^6 particles/cm³). Particle size distribution indicated the geometric mean diameter of the generated particles to be average 15 nm (Cu), 25 (Zn) and 16 nm (welding rods) with a geometric standard deviation of 1.5, 1.7 and 1.4, respectively. SEM and TEM results confirmed the deposition of nano-sized particles on cell-free transwells. The cellular concentration of Cu NPs was 5.9 μg Cu/transwell (4.7 cm²) and a substantial amount of Cu was released to the basolateral medium (0.3 μg) during air-delivery of Cu NPs in 4 h. Viability for cells exposed to Cu NPs was significantly reduced to 79% (p<0.05) at 4 h post-exposure compared to cells maintained in an incubator. Our results demonstrated that the SDS can be useful for generating metal NPs and simulating welding fumes for their toxicity assessment. In addition, integrated ALI exposures of human lung cells with SDS can provide screening data to aid prediction of toxicity of metal NPs.

Engineered nanoparticles are being used in various commercial products; however, a significant concern is the potential of these particles to cause adverse effects on human health, as evidenced by exacerbation of allergic airway disease. The objective of this interdisciplinary research is to investigate the putative adjuvant properties of engineered nanoparticles on biological responses at the animal, cell, and membrane level. Two types of nanoparticles (SNPs) with different size and surface coating were modified with alkyne-terminated surfaces and appended with polyethylene glycol azides via ‘click’ chemistry. At the cell level, in vitro mouse models were utilized to examine the effect of modified SNPs on dendritic cell (DC)-induced T cell effector function. Ovalbumin (OVA)-(derived peptide OVA57-65, or OVA323-339 were presented by endogenous antigen presenting cells in LPS or a mouse DC line DCS. In an in vivo setting, intratracheal injection of CD8 or CD4 T cells in response to OVA peptides was measured by flow cytometry. At modest peptide stimulation levels, modified SNPs (up to 10 μg/ml) enhanced the proportion of CD8, but not CD4, T cells that produced cytokines. Various functional groups ((-COOH, -NH₂, -OH) on modified SNPs enhanced cytokine production to different levels, with -COOH SNPs being the most effective. Furthermore, 50 nm -COOH SNPs exhibited greater enhancement effect on CD8 T cell response than the other sizes. Importantly, modified SNPs did not aggregate in in vitro culture media, demonstrating their effect at the true nanoscale. Collectively, our results demonstrated the potential adjuvant effect of modified SNPs on CD8 T cell function and will also complement other studies being conducted by the team at the membrane and the intact animal level. (Supported by NIH Grant RC2 ES018756)

The risk posed by lead during manufacturing of ferroelectric ceramics has led to efforts to develop lead-free options, bismuth sodium titaneum barium titaneum (BNT-BT) nanoparticles (NP) have become a strong candidate to replace lead zirconate titanate (PZT)-based ceramics. In order to evaluate the biological effects generated by PZT and BNT-BT NP we investigated their interaction with biomolecules adsorbed on the NP surface, which defines the interface with cellular membranes, the uptake and potential toxicity in human cell lines. The NP were dispersed, stabilized with bovine serum albumin (BSA) and characterized in complete cell culture medium showing an average size of 188 nm (BNT-BT) and 168 nm (PZT), as measured by differential centrifugal sedimentation. The analysis of the proteins adsorbed with high affinity to the NP surface (hard corona) after the incubation with FBS showed prevalently BSA and complex protein profiles dependent on the concentration of FBS. In vitro uptake studies in cell cultures rely on fluorescently labeled NPs; we generated BNT-BT and PZT NP showing fluorescent properties by labeling BSA used to stabilize the dispersions. The NP were tracked through the endosomal pathway using a time lapse live cell imaging approach. For cytotoxicity assessment, cell membrane damage and mitochondrial activity were measured in Hep G2, LLC-PK1, A549 and SH-SYSY cells after exposure to increasing concentration of BNT-BT and PZT NP for 26 or 48 h. The integrity of the cell membrane was not significantly affected; however the mitochondrial activity increased after 48 h with 100 μg/ml for both NP being more evident for the PZT NP in neutrophiloma cells. Our results suggest that BNT-BT showed lower levels of toxicity compared to PZT thus it could be a good replacement for BNT-based ceramics. Funding from the European Community Seven Framework Program and CONA-CVT (Grant agreements 262387 and 12514) and Epitope Map Program.

Airborne nanoparticles (NPs) that reach the alveolar region are likely to be presented to alveolar epithelial cells coated with a corona of lung surfactant lipids and proteins. However, the effect of the core material, airborne NP toxicity is still unclear, mainly because most studies were conducted in cells exposed to NPs in growth media, limiting the corona to molecules found in the media. Furthermore, accumulating observations support a role for positive surface charge in NP toxicity, but the impact of the corona, which shifts the NP surface charge to more negative values, is unclear. To understand the role of corona in airborne NP exposure, we exposed alveolar epithelial cells at the air-liquid interface (ALI) to aerosolized NPs that were pre-incubated with a natural surfactant lipid and protein (Infasurf®) or BSA solution. Bare polystyrene (PS) NPs, showing no toxicity, were compared with aminated or carboxylated PS NPs, showing high or low level of toxicity, respectively. Cellular response was compared between NPs at the same cellular dose, measured as μg/cm² using a quartz crystal microbalance. We found that the lowest dose required to induce toxicity by aminated NPs with no corona was 3 fold lower than the dose required to induce toxicity by aminated NPs with a surfactant co-
rona, and 1.8 folds lower than the dose required by aminated NPs with a BSA corona. In contrast, BSA corona eliminated toxicity of carbonylated NPs at all measured doses. With no corona, the level of aminated NPs at the cell surface or inside the cytoplasm was significantly higher than the level of carbonylated NPs. To evaluate the possibility that the observed protective effect of surfactant or BSA coronas from aminated or carbonylated NP toxicity, respectively, is due in part to a decrease in NP-cell interaction and internalization, we currently quantify any changes in these processes that might be induced by the coronas. This work was funded by NIEHS grant 1R22ES018786-01 to GO.

1763 Metastatic Capacity Induced by Cells Exposed to Titanium Dioxide Nanoparticles.
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Background. Titanium dioxide nanoparticles (TiO2 NPs) have been classified as a possibly carcinogenic to humans (Group 2B) by the International Agency for Research in Cancer. In this regard, its potential carcinogenic effects have been under research. However, less has been investigated about the effects of exposure to TiO2 NPs when they reach bloodstream, especially if TiO2 NPs exposed cells could proceed from a tumor. Aim. We hypothesized that tumor cells exposed to TiO2 NPs could promote a metastatic events. To test this hypothesis, lung adenocarcinoma cells were exposed to TiO2 NPs (1, 5 and 10 μg/cm2) for 7 days and 30,000 cells of each concentration were injected into bloodstream of chicken chorioallantoic membrane (CAM) of free pathogens fertilized eggs previously incubated during 11 days at 37°C and 80% humidity. Analysis of morphological changes was performed at 16th day. Results. TiO2 NPs were characterized by transmission electronic microscopy (TEM) suspended in F12K medium with 10% Serum Fetal Bovine (SFB) and zeta potential was -20.98±0.75, and TiO2 NPs agglomerate size was 212.3±4.417nm, 630.2±20.94 nm and 588.3±630.2 nm for 1, 5 and 10 μg TiO2/mL respectively. Images showed that there is an increase in distance between blood vessels and a decrease in number of nodes in CAM treated with cells exposed to 10 μg/cm2 of TiO2. In addition, there were observed cell clusters on the CAM treated with cells exposed to 1 and 5 μg/cm2 of TiO2 NPs. Conclusion. TiO2 NPs exposure (10 μg/cm2) to lung adenocarcinoma cells induced an increase in blood vessel distance and a decrease the number of nodes, which seems to be a downregulation effect in angiogenesis. However, pre-exposed lung adenocarcinoma cells to 1 and 5 μg/cm2 of TiO2 NPs provided a capacity to form clusters, which could represent a higher metastatic effect.

1764 Phenotypic Polarization and Attenuation of Toll Receptor Signaling Functions in Macrophages Exposed to Engineered Nanoparticles.
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Although macrophages play a critical role in scavenging engineered nanoparticles (ENPs) from tissue, relatively little is known about how their normal innate immune function is impacted. We investigated how pretreatment of macrophages with 33 nm superparamagnetic iron oxide (SPIO) or 50 nm amorphous silica modulated the transcriptional response of macrophages to the toll 4 receptor (TLR4) ligand LPS. Affymetrix microarray analysis showed over 5000 mRNA's were differentially regulated (≥1.5 fold, p<0.05) in response to LPS (10 ng/ml, 6 hrs). SPIO pretreatment (25 μg/ml) alone altered expression of a smaller set of genes (1029 total), but modulated expression of nearly 560 LPS regulated genes in a greater than additive manner. In contrast to SPIO, relatively few LPS regulated genes were modulated by silica pretreatment. Pathway analysis showed that pretreatment with SPIO suppressed LPS activation of several key cellular functions like chemotaxis, interferon and Jak/Stat inflammation signaling while enhancing cell adhesion and oxidative/nitrative stress responses. Pretreatment of macrophages with SPIO also caused a dose dependent decrease in phagocytosis of S.pneumoniae and S typhimurium, whereas silica pretreatment had no effect. Flow cytometry studies using fluorescent LPS or antibodies against TLR4 revealed that SPIO exposure caused dose-dependent down regulation of cell surface TLR4 level while silica had no effect. Our results demonstrate that macrophages exposed to SPIO NPs are polarized toward an anti-inflammatory phenotype which attenuates pro-inflammatory signaling by TLR4, and suppresses their phagocytic activity against pathogens.

This work was supported by NIEHS Grant U19-ES019544.

1765 Intracellular Trafficking and Accumulation Dynamics of Zinc Ions in Alveolar Epithelial Cells Exposed to Airborne ZnO Nanoparticles at the Air-Liquid Interface.

Airborne nanoparticles (NPs) that enter the respiratory tract are likely to reach the alveolar region. Accumulating observations support a role for zinc oxide (ZnO) NP dissolution in toxicity, but the majority of the studies were done in cells exposed to NPs in growth media, where large doses of dissolved ions are shed into the exposure solution. To better represent the exposure in the respiratory tract and focus on the dissolution of airborne NPs in the cellular environment, we exposed alveolar epithelial cells to aerosolized NPs at the air-liquid interface (ALI). Using a fluorescent indicator for zinc ions (Zn2+) and organelle-specific fluorescent proteins, we quantified Zn2+ in single cells and organelles over time and correlated these values with toxicity. We found that intracellular Zn2+ in cells exposed at the ALI peaked 3 h post exposure and decayed to normal levels by 24 h, which was in contrast to exposure in submerged cultures where intracellular Zn2+ continued to increase over time. For the lowest toxic dose at 24 h, the peak at the ALI was ~3 folds lower than the 24 h value in submersed cells, and ~8 folds lower than the 24 h value in submersed cells exposed to Zn2+ only. At the ALI, 45% of intracellular Zn2+ was found in endosomes at 1 h post exposure, decreasing to 24% by 3 h. In contrast, 20% of intracellular Zn2+ was found in lysosomes at 1 h, increasing to 42% by 3 h. Interestingly, exposure of submersed cells to Zn2+ only, led to a minimal accumulation of ions in either the endosomes or lysosomes, with the majority of ions found in larger vesicles. Our observations indicate that airborne ZnO NPs induce toxicity at the ALI at intracellular Zn2+ levels that are significantly lower than those detected when toxicity is induced in submersed cultures. The localized dissolution and trafficking of Zn2+ in endosomes following with their accumulation in lysosomes play critical roles in airborne NP toxicity. This work was funded by NIEHS grant 1R22ES018786-01 to GO.

1766 Aggregation Dynamics and Structure Measurements of Nanomaterials in Biologically Relevant Conditions.
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Nanomaterial dosimetry for in vitro studies continued to be debated in nanotoxicology literature, particularly due to the propensity of aggregation of highly diffusive nano-scale materials. In classical nanotoxicology literature, average particle size and surface charge are the typical measured parameters either initially or at the end of exposure time period; with no information on the dynamic behavior throughout the process. Moreover, aggregate structural information is mostly ignored. This study focuses on monitoring dynamic aggregation behavior of metallic and carbonaceous nanoparticles under biological (i.e., in exposure media with added penicillin streptomycin, at 37 °C) exposure conditions. The aggregation dynamics is monitored by employing state-of-the-art ALV goniometer system. Suspension phase fractal dimension was measured with angle-dependent static light scattering for an angular range of 12°-120°. Aggregation dynamics results indicate that electrostatic interaction serves as the key interfacial force to providing stability to nanomaterials. In addition, continuous measurements of aggregate size in physiological condition show network formation. Furthermore, time dependent fractal dimension results indicate that aggregate structure remains unchanged over time; however, significantly altered by the background solution chemistry. The key implications of the study include: consideration of intermediate time-endpoint (instead of 24 h); aggregation dynamics and network formation can minimize size effects on nanotoxicology; aggregation dynamics with aggregate structure information can better determine nanomaterial dosimetry.

1767 Targeting Alternatively Activated Rat Macrophages Using Mannose Functionalized Nanocarriers.
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In response to inflammatory signals generated at sites of injury, macrophages are activated into two main phenotypes: classically activated M1 cells and alternatively activated M2 cells. Whereas M1 macrophages release cytotoxic/proinflammatory
mediators that contribute to tissue injury. M2 macrophage-derived mediators suppress inflammation and initiate wound repair. Mannose receptor (MR) is known to be expressed at high levels on M2 macrophages. In these studies, a mannose functionalized nanocarrier (NC) was designed and developed with the goal of selectively targeting M2 macrophages. Mannose functionalized NC (FTTC-Gaba-Ser-Man)-PEG12-Ser-Man)-Gaba-Gaba-Cys) were prepared using solid phase synthesis and their identity confirmed by MALDI-TOFMS and HPLC analysis. Rat peritoneal macrophages (PMs) were incubated for 24–48 h with IFN-γ (20ng/ml) or IL-4/IL-13 (10ng/ml), to induce M1 or M2 activation, respectively. mRNA levels of the M1 marker, inducible nitric oxide synthase (iNOS) and the M2 marker, arginase (Arg)-1 were quantified by RT-PCR, and protein levels by western blotting. Treatment of M0 with IFN-γ resulted in increased expression of iNOS mRNA and protein for 60 min, NC binding to macrophages was quantified by confocal microscopy. Confocal microscopy showed that uptake of mannosylated NC into M2 PMs was 2.4 fold greater than control PMs, and 1.8 fold greater than M1 PMs. M2 macrophage phenotype-specific uptake was completely abolished by mannan, suggesting MR-mediated targeting. NCs were co-localized with rhodamine-dextran, a marker for cells of macrophage phenotype. Experiments showed that uptake of mannosylated NC into M2 PMs was 86.6% while Bi2O3 and ZV-Bi in 27.8 and 26.1%, respectively. Uptake was assessed in cells of various types, including lung cells, and was found to be higher in lung cells compared to other cell types.

Therefore, NP transport and uptake is a relevant process that needs to be thoroughly investigated. Testing for biocompatibility and toxicity is essential to ensure that nanomaterials are safe for use in medical applications. We investigated physicochemical properties of nano-sized oxides of Fourth Period transition metals that contribute to cytotoxicity. The cytotoxicity (i.e., cell killing) of nanoparticles (NPs) TiO2, Cr2O3, Mn2O3, Fe2O3, NiO, CuO, and ZrO2 increases as atomic number increases. This trend is not cell type specific, as it is observed in nontransformed human lung (BEAS-2B) and adenocarcinomic human alveolar basal (A549) epithelial cell lines. We assessed physicochemical properties of NPs to discover the determinants of cytotoxicity: 1) the point-of-zero charge (PZC) (i.e., isoelectric point) describes the surface charge of NPs in cytosolic and lysosomal compartments; 2) the relative number of available binding sites on the NP surface; and 3) the relative amount of available binding sites on the NP surface.

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RESULTS: Electronic microscopy studies revealed an average Np size of 130 nm (σ = ±32.7 nm) and irregular spherical in shape. IR, EDS and DLS results support the formation of NPs detecting mineral and organics constituents respectively with high stability. The in vitro tests shows by first time the Nps-α-TOS can be internalized and are more cytotoxic and effective that α-TOS alone and inhibits the growth of resistant cervix cancer cell.

CONCLUSION: In this study we found that magnetite Nps can work as nanocarriers of α-TOS. This composite protect the anticancer activity of α-TOS and enhances the anti-tumor effect in resistant cancer cells.

Engineered nanoparticles (NPs) possess numerous potential benefits to society in fields as diverse as electronics, textiles, medicine, energy and construction. The respiratory system represents a unique target for the potential toxicity of NPs. Due to their dimensions, inhaled NPs can reasonably be expected to penetrate to the deepest part of the lungs, the alveolar sacs. Completed inhalation studies in laboratory rats have demonstrated that some NPs induce oxidative stress, inflammation and fibrosis. In this study, we examine the effects of nanomaterial and parent nano-materials (bismuth oxide [Bi2O3] and copper oxide [CuO] spheres and rods) on an alveolar co-culture model consisting of human type II pneumocytes and alveolar macrophages. Co-cultures were set up with a ratio of 3:1 of type II pneumocytes to each alveolar macrophage 24 h prior to treatment. Media was then removed and co-cultures were treated with 0.001 100 mg/ml NPs for 24 h. Cell viability was only affected by CuO at 100 mg/ml; CuO spheres and rods had similar effects. These studies demonstrate that only high concentrations of raw Bi2O3 and CuO NPs affect respiratory co-cultures and that CuO NP size does not affect respiratory co-cultures. Additional phagocytosis studies and cytokine analysis studies (intracellular and secreted concentrations) are being conducted to evaluate inflammatory effects of these NPs in the respiratory co-culture immune cells.

The cellular internalization mechanism of engineered nanoparticles (NPs) and their intracellular trafficking govern the cellular interactions of the particles, which ultimately determine their impact on the cell. These cellular mechanisms largely depend on the size of the NPs or the aggregates that are often formed. To identify mechanisms specific to individual NPs or small nano-scale aggregates, we used stoichiometric optical reconstruction microscopy (STORM) to resolve and accurately localize the position of individual NPs within organelles and in respect to subcellular structures with 10-20 nm resolution. Using FastLime, a photo-switching derivative of green fluorescent protein, we created fluorescent chimeras for clathrin, caveolin and actin, and expressed the proteins in alveolar epithelial cells. The transfected cells were exposed to amorphous silica NPs, tagged with a photo-switching dye, and imaged at 2 h and 16 h post exposure. We found NPs within clathrin-coated pits and caveolin-coated vesicles in both time-points, suggesting a preferential binding of the NPs to molecules found in clathrin-coated pits and caveolae at the cell surface. A significant number of NPs were found aligned along actin filaments at 16 h post exposure. The distance between the NPs and the filaments was calculated for more than 1100 NPs, and the distribution of the distances in the range of 0 to 100 nm was significantly different from the distribution expected to occur if the NPs were randomly placed in the cytoplasm. Together, our observations suggest a mechanism for the internalization and trafficking of amorphous silica NPs in alveolar epithelial cells that occurs through interactions with molecules found in caveolae and clathrin-coated pits, followed by internalization via vesicles that eventually are shuttled along actin filaments within the cytoplasm. This work was funded by NIEHS grant 1RC2ES018786-01 to GO.

The effect of titanium dioxide nanoparticles (nano-TiO2, Degussa p25, 86% anatase and 14% rutile) treatment of human lung epithelial cells (BEAS-2B) was examined by analyzing changes in messenger [mRNA] and microRNA [miRNA]. BEAS-2B cells were treated with 0, 3, 10, 30 or 100 ug/ml nano-TiO2 for 1 day (for mRNA analysis) or 3 days (for miRNA analysis). Differentially expressed mRNA and miRNA were analyzed using Affymetrix microarrays (human U133 Plus 2.0) and Affymetrix microarrays, respectively. Although, the tested doses were not cytotoxic, there were alterations in both mRNA and miRNA expression. The expression of mRNA/miRNA changes were examined in MetaCore (GeneGo) and IPA (Ingenuity Pathway Analysis) to delineate associated signaling pathways. Signaling pathways altered by nano-TiO2 treatments included cell cycle regulation, apoptosis, calcium signaling, translation, NRF2 – mediated oxidative response, IGFl signaling, RAS signaling, PI3K/AKT signaling, cytoketone remodeling, cell adhesion, BMP signaling, and inflammatory response. Many of the genes in these signaling pathways are known to be regulated by the miRNAs that were altered by the nano-TiO2 treatment. The mRNA 17-92 cluster and let-7 miRNA family that are reportedly involved in lung cancer formation were altered by nano-TiO2 treatment. The miR-17-92 cluster, an oncogenic microRNA cluster, is induced while the tumor suppressor microRNA, let-7 family, is suppressed. The observed changes in miRNA expression introduces an additional mechanistic dimension that supports the significance of the observed miRNA expression changes, and demonstrated that the nano-TiO2 treatment can cause diverse but coordinated pathway alterations associated with changes in vivo response to tumorigenesis. [This abstract does not necessarily reflect the policies of the U.S. EPA.]
and abluminal sides, respectively, of a transwell membrane. Permeability studies using fluorescent carboxyfluorescein tagged extracellular polypeptide (PEG) confirmed a tight layer of bEnd3 cells with transendothelial resistance of 122 ± 3 Ωcm2. This culture set up was treated with citrate and PEG-coated Au-NPs (50 μg/ml) for 24 hr to quantify non-specific leakage of substrates across the cell barrier and to evaluate the association of NPs with astrocytes, with and without flow of media. Immediately coupled plasma mass spectrometry results show fewer NPs on the abluminal side confirming a tight cell barrier and transmission electron microscopy imaging suggest endosomal localization of NPs in bEnd3 cells. Unlike the static set up, brightfield imaging suggested less aggregation of NPs with flow and citrate-coated NPs associate more with astrocytes in comparison to PEGylated NPs. These data demonstrate that surface properties influence passage of NPs across bEnd3 cells. This multiregional model could be applied as a rapid screening technique to analyze uptake and toxicity of NMs across the BBB.

### 1777 Effect of Surface Modification of Metal Oxide Nanoparticles upon Cell Viability and Genotoxicity of Epithelial Breast Cells


In this work, the biological activity of different metal oxide nanoparticles (Fe3O4, Co3O4, CuO) upon epithelial breast cells MCF-10A have been evaluated. The first step is the fabrication and characterization of the metal-oxide nanoparticles (MONPs) followed by the functionalization with folic acid, L-arginine and L-cysteine which is confirmed by Fourier Transform Infrared Spectroscopy (FTIR) and Low Voltage Electron Microscopy (LTEM) measurements. In order to determine real time cell viability, semi-confluent cells were exposed to either 0 or 0.2 mg/mL of particles with and without functionalization. 2 x 103 cells/well were grown in a gold wired 16-well plate for 15 hours, with readings of impedance collected every 15 minutes to monitor cell proliferation. To determine cell-nanoparticle interactions, cells were grown for 24 hours, exposed to 2 mg/ml of nanoparticles (PEG) lowered to grow for additional 48 hours. After that, cells were fixed and stained with DAPI to evaluate apoptosis necrosis and characterization of normal nuclear morphology. Additionally a dose-response growth curve using Fe3O4 nanoparticles, presumably, the less toxic exposure, free and functionalized with folic acid (Fe-AF) was carried out. Results indicate that the most toxic treatment was Fe3O4 and the less toxic was Fe3O4 functionalized with folic acid, although all the MONPs evaluated didn't cause a strong toxic response due to the values of cell index obtained by impedance evaluation. However, the presence of particles interacting with the cell and nucleus showed the possibility of genotoxic effects, and for that reason, in an attempt to explore this condition, the micronucleus assay employing Cytchalasin B (Cytome assay) is being conducted as a follow up.

### 1778 Synthesis, Characterization and Evaluation of Biological In Vitro Activity of Eu3+ Doped Hydroxyapatite


In this work, the synthesis, characterization and biological in vitro activity of Eu3+ doped hydroxyapatite (Eu-HAp) is being evaluated. Hydroxyapatite synthesis is being carried out by microwave-hydrothermal techniques, modifying parameters such as temperature, power, pH and calcium precursors. In a typical process, the synthesis was carried out by the mixture of CaCl2 or Ca(NO3)2·4H2O and Eu(NO3)3·5H2O. After that, a solution of (NH4)2HPO4 was added. The full synthesis was carried out by the mixture of CaCl2 or Ca(NO3)2·4H2O and Eu(NO3)3·5H2O. After that, a solution of (NH4)2HPO4 was added. The full

### 1779 Interactive Toxicity of Uronic Acid and Lipopolysaccharide in Human Liver HepG2 Cells


Uronic acid (UA), a natural botanical product, is a constituent of some dietary supplements used for weight loss. It has been associated with clinical hepatotoxicity leading to liver failure in humans. The present study was undertaken to evaluate the interactive toxicity, if any, of UA with lipopolysaccharides (LPS), a potential contaminant of food, at low nontoxic concentrations. The human hepatoblastoma HepG2 cells were treated with the vehicle control and test agents, separately and in combination (UA+LPS) at concentrations of UA 1.0 μM and LPS 1.0 ng/ml, for 24 h at 370 C in 5% CO2. Following the treatment period, the cells were evaluated by the traditional biochemical endpoints of toxicity in combination with the toxicogenomic endpoints that included cytotoxicity, oxidative stress, mitochondrial injury and changes in pathway-focused gene expression profiles. Compared to the controls, low nontoxic concentrations of UA and LPS separately showed no effect on the cells as determined by the biochemical endpoints. However, the simultaneous mixed exposure of the cells to their mixture resulted in increased cytotoxicity, oxidative stress and mitochondrial injury. The pathway-focused gene expression analysis resulted in the altered expression of several genes out of 84 genes examined. Most altered gene expressions induced by the mixture of UA and LPS were different from those induced by the individual constituents. The genes affected by the mixture were not modulated by either UA or LPS. The results of this study suggest that the interactions of low nontoxic concentrations of UA and LPS produce toxicity in HepG2 cells.

### 1780 Dihydroartemisinin Inhibits PMA-Induced Cyclooxygenase-2 Expression through Downregulating AKT and MAPK Signaling Pathways in Murine Macrophages

E. Han1, H. Kim1, J. Choi1, Y. Hwang1 and H. Jeong1

Dihydroartemisinin (DHA), a semi-synthetic derivative of artemisinin isolated from the traditional Chinese herb Artemisia annua L., has recently been shown to possess antitumor activity in various cancer cells. However, the effect of anti-inflammatory potentials of DHA in murine macrophage RAW 264.7 cells has not been studied. The present study investigated the effect of COX-2 and molecular mechanisms by DHA in phorbol 12-myristate 13-acetate (PMA)-stimulated RAW 264.7 cells. DHA dose-dependently decreased PMA-induced COX-2 expression and PGE2 production, as well as COX-2 promoter-driven luciferase activity. Additionally, DHA decreased luciferase activity of COX-2 regulation-related transcription factors including NF-kB, AP-1, C/EBP and CREB. DHA also remarkably reduced PMA-induced p65, C/EBPβ, c-jun and CREB nuclear translocation. Furthermore, DHA evidently inhibited PMA-induced phosphorylation of AKT and the MAP Kinases, such as ERK, JNK and p38. These data indicated that DHA efficaciously attenuates COX-2 production via down-regulation of AKT and MAPK pathway, revealing partial molecular basis for the anti-inflammatory properties of DHA. These findings demonstrate that DHA effectively attenuates COX-2 production, and provide further insight into the signal transduction pathways involved in the anti-inflammatory effects of DHA.

### 1781 Protective Role of Silymarin against Oxidative Stress Induced in Human Neuroblastoma Cell Line SH-SY5Y

A. Anadon, M. A. Martinez, E. Ramos, V. Castellano, I. Ares, M. Martinez, M. R. Martinez-Larranaga and A. Romero

Silymarin (SM) is a mixture of bioactive flavonolignans isolated from Silybum marianum (L.) Gaertn., employed usually in the treatment of alcoholic liver disease and as anti-hepatotoxic agent in humans. The essential activity of SM is an antioxidant activity of Eu3+ Doped Hydroxyapatite.

In this study, the mixture of UA and LPS was confirmed in the samples obtained by the most alkaline pH ranges (above 10), while at pH 9 we obtained a mixture between hydroxyapatite and monetite phases. To evaluate the biological in vitro activity of our materials, MTT assay is being performed upon mouse fibroblasts that are treated with representative samples of Eu-HAp nanoparticles previously synthesized by our group.
effect of its flavonolignans. Because of the importance of oxidative stress and mito-
condrial dysfunction in causing neuronal death, prompted us to investigate the ef-
effects of SM against an in vitro model of reactive oxygen species (ROS) production
in the human neuroblastoma cell line SH-SY5Y. We selected two cytotoxic stimuli,
for one hand, hydrogen peroxide (H2O2) (500 μM), and on the other hand the
combination of 30 μM rotenone plus 10 μM oligomycin-A (R/O) that inhibit mi-
tochondrial respiration complexes I and V, respectively. Cell viability, measured as
MTT reduction, was decreased to 70% in cells treated with H2O2 and to 60% in
cells exposed to R/O. Cell incubation with increasing concentrations of SM (1-
1000 μM) for 24 hr, followed by a 24-hr period with H2O2 (extracellular ROS) or
R/O (intracellular ROS). Maximum protection was achieved with 300 μM SM
(30% protection). Our results showed that R/O and H2O2-induced cytotoxicity in
SH-SY5Y cells was suppressed by treatment with SM. Because, it is recently re-
ported that SM crosses the blood–brain barrier and enters the CNS and it is non-
toxic even at higher doses, this flavonoid may be useful in diseases known to be ag-
gravated by reactive oxygen species and in the development of novel treatments for
neurodegenerative disorders. This work was supported by projects Ref. BSCHGR85/08(UCM), Ref. No. S2009/AGR-1469(CAM) and Consolider-
Ingenio 2010 No.CSD2007-063(MEC), Spain.

1782 Thymoquinone, a Bioactive Component of Nigella sativa, 
Modulates Redox Status and Insulin Secretion from 
Pancreatic Beta Cells.
J. P. Gray 1, 3, R. Follmer 1, R. Rebar 1, N. Seeram 2 and E. Heartl 1.
1Science, US Coast Guard Academy, New London, CT; 2Bioactive Botanical Research Laboratory, University of Rhode Island, Kingston, RI; 3Cellular Dynamics, Marine Biological Laboratory, Woods Hole, MA.
Nigella Sativa is a traditional medicine that has been used in the Mediterranean to treat a variety of disorders, including type 2 diabetes. A primary component of Nigella sativa extract is thymoquinone which, like Nigella sativa extract, attenuates diabetes symptoms. The molecular targets and interactions of thymoquinone with metabolic pathways relevant to glucose-stimulated insulin secretion (GSIS) from pancreatic β-cells have not yet been identified. Our laboratory previously demonstrated that low (nM-μM) doses of various quinones such as menadione stimulate insulin secretion from β-cells, and this action was coupled to the generation of low levels of H2O2, a putative mediator of GSIS. Here we compared the mechanism of action of thymoquinone to that of menadione in β-cells. Like menadione, thymo-
quinone induced a dose-dependent increase in the production of H2O2. Unlike menadione, the redox cycling of thymoquinone was not dependent on the glucose concentration. Both NADPH and NADH supported the redox cycling of thymo-
quinone in cytosolic and mitochondrial fractions. This was consistent with the abil-
ity of thymoquinone to decrease NADH/NAD+ and NADPH/NADP+ ratios, thus
reducing intracellular redox poise. Thymoquinone-dependent redox cycling activi-
ties were inhibited by diphenylene iodonium, an inhibitor of flavin-containing ox-
idoreductases. Dicoumarol and MAC220, NQO1 inhibitors previously shown to in-
hbit menadione-dependent redox cycling, failed to inhibit thymoquinone-de-
pendent redox cycling. Unlike menadione, thymoquinone was found to potentiate GSIS in a dose-dependent manner at stimulatory glucose (concentrations which potently stimulate insulin secretion). These data suggest that while the mechanisms of thymoquinone redox cycling are different than those of menadione, thymo-
quinone retains the ability to regulate both redox status and insulin secretion from
β-cells.

Antioxidant and DPPH-Scavenging Activities of Compounds and 
Ethanolic Extract of the Leaf and Twigs of Caesalpinia 
bonduc L. Roxb.
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A. A. Akindahunsi 1 and N. H. Tan 1.
1Biological Sciences, Covenant University, Ota, Nigeria; 2Biological Sciences, Cranfield University, Old, Nigeria; 3Medical Biochemistry, College of Medicine, Lagos State University, Ikeja, Nigeria; 4Biochemistry, Federal University of Technology, Akure, Nigeria; 5State Key Laboratory of Physicochemistry and Plant Resources, Kunming Institute of Botany, Kunming, China.
Antioxidant effects of ethanolic extract of Caesalpinia bonduc and its isolated bioac-
tive compounds were evaluated in vitro. The compounds included two new cascade
diterpenes, 1,7α-diacetoxy-5,6-dihydroxy-cass-14(15)-epoxy-16,17-olide (1) and
12α-ethoxy-1,14β-diacetoxy-2,5α-dihydroxycass-13(15)-en-16,12-olide (2); and others, bonducullin (3), 7,4′-dihydroxy-3,11-dehydrohomoisolavone
(4), daucosterol (5), luteolin (6), quercetin-3-methyl ether (7) and kaempferol-3-
O-β-D-glucopyranoside (8). The antioxidant proper-
ties of the extract and compounds were assessed by the measurement of the total
phenolic content, ascorbic acid content, total antioxidant capacity and 1,1-
diphenyl-2-picryl hydrazyl (DPPH) and hydrogen peroxide radicals scavenging ac-
tivities. Compounds 3, 6, 7 and ethanolic extract had DPPH scavenging activities
with IC50 values of 186, 75, 17 and 102 μg/ml respectively when compared to vita-
min C with 15 μg/ml. On the other hand, no significant results were obtained for
hydrogen peroxide radical. In addition, compound 7 has the highest phenolic con-
tent of 0.81±0.01 mg/ml of gallic acid equivalent while compound 8 showed the
highest total antioxidant capacity with 254.3±1±3.45 and 199.8±2±7.8 μg/mg gallic
and ascorbic acid equivalent respectively. Compound 4 and ethanolic extract
showed a high ascorbic acid content of 2.26±0.01 and 6.78±0.03 mg/ml respec-
tively. The results obtained showed the antioxidant activity of the ethanolic extract of C. bonduc and deduced that this activity was mediated by its isolated bioactive compounds.

3-Caffeoyl, 4-Dihydrocaffeoylquinic Acid from Salicornia 
herbacea Attenuates High Glucose-Induced Hepatic 
Lipogenesis in Human HepG2 Cells.
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3-Caffeoyl, 4-dihydrocaffeoylquinic acid (CDQC) from Salicornia herbacea has a variety of pharmacological properties, including antioxidant and anti-inflammatory and hepato-protective properties. The aims of our study were to provide new data on the molecular mechanisms underlying the role of CDQC in prevention of high glucose-induced lipid accumulation in human HepG2 cells. We found that CDQC suppressed high glucose-induced lipid accumulation in HepG2 cells. CDQC strongly inhibited the high glucose-induced FAS expression by modulating SREBP-1c activation. Moreover, the use of a specific inhibitor or liver kinase B1 (LKB1)-sirtuin A transfected HepG2 cells showed that CDQC activated AMP-acti-
vated protein kinase (AMPK) via silent information regulator T1 (SIRT1) or LKB1 in HepG2 cells. These results indicate that CDQC prevents lipid accumulation by

1784 Selective Elimination of Malignant Melanomas through 
Autophagic and Mitochondria-Based Mechanisms by the 
Antitumor Agent OsW-1.
K. Riaz Ahmed, C. Garcia-Prieto, L. Feng and P. Huang. Molecular Pathology, University of Texas MD Anderson Cancer Center, Houston, TX; Sponsor: M. Smith.
Drug resistance and lack of therapeutic selectivity are two of the biggest challenges to successful melanoma therapy. Constitutive activation of Extracellular Signal Regulated Kinase 1/2 (ERK1/2) and subsequent chemo-resistance has been reported in malignant melanomas. ERK1/2 has also been implicated in activation of mito-
chondrial gene expression and regulation of autophagy thus making it an important therapeutic target.

The natural product, OSW-1, isolated from the bulbs of ivory coast lily, has been shown to be highly cytotoxic in numerous cancer cell lines with yet undefined mechanisms of action. Herein, we report our results on the anticancer activity and selectivity of OSW-1 in malignant melanoma cells and its potential mechanisms of action.

Our preliminary results demonstrated that OSW-1 was highly effective in killing tumor cells that are resistant to most of the currently available anticaner drugs, with IC50 values in sub-nM concentrations. Importantly, OSW-1 preferentially killed melanoma cells and exerted much lower toxicity to normal melanocytes in culture. Biochemical analysis revealed that OSW-1 treatment caused damage of the mitochondrial membrane integrity, leading to a decrease in transmembrane potent-
tial and subsequently initiating cell death, apparently through autophagy. Further study demonstrated that OSW-1 inhibited ERK1/2 expression in melanoma cells and caused a significant disturbance of cellular calcium homeostasis, leading to aberrant calcium-mediated processes including mitochondrial impairment. Based on these results, we postulate that OSW-1 inhibits ERK1/2 mediated signaling and triggers mitochondrial damage in cells leading to a significant disturbance of cellular calcium and cell death through autophagy. This study is of great significance since ERK1/2 signaling is involved in melanoma survival and inhibition of ERK1/2 expression and induction of autophagic cell death by OSW-1 will be criti-
cal to combat therapeutic resistance and enhance drug selectivity.
Epithelial to mesenchymal transition (EMT) is a key event in the progression of cancer. EMT is characterized by the loss of epithelial and the gain of mesenchymal features. Previous studies have revealed that treatment with CKS, saponins from the roots of Platycodon grandiflorum, significantly reduces metastasis and tumorigenesis, but the underlying mode of action has not been elucidated. In this study, we investigated the inhibitory effect of CKS on transforming growth factor (TGF)-β1-induced alterations characteristic of EMT in human lung carcinoma cells. We found that CKS-treated cells displayed inhibited TGF-β1-mediated E-cadherin down-regulation and Vimentin up-regulation and also retained epithelial morphology. Furthermore, TGF-β1-induced Snail expression was reduced by CKS. Pretreatment of cells with CKS blocked TGF-β1-induced Smad2/3 phosphorylation and Smad7 down-regulation. CKS inhibited TGF-β1-induced phosphorylation of Akt, ERK1/2, and glycogen synthase kinase-3β (GSK-3β). Furthermore, TGF-β1-induced Snail expression was reduced by pharmacology inhibitors of Akt, ERK1/2, and GSK-3β. These results indicate that pretreatment with CKS inhibits the TGF-β1-induced EMT process and prevents TGF-β1-induced transdifferentiation via activation of Akt and ERK1/2 and inactivation of GSK-3β in A549 cells.

Platycodin D, the saponins from the roots of Platycodon grandiflorum (CKS), has a variety of pharmacological properties, including anti-hyperlipidemic, antioxidant and hepatoprotective properties. This study was conducted to suggest the role of AMP-activated protein kinase (AMPK) and inhibited NF-κB as well as CREB activation.

Effects of Saponins from the Roots of Platycodon grandiflorum on TGF-β1-Induced Epithelial-Mesenchymal Transition in in A549 Cells.

C. Ho1, Y. Hwang1, Y. Chung2 and H. Jeong1, 1Pharmacy, Chungnam National University, Daejeon, Republic of Korea; 2International University of Korea, into, Republic of Korea.

Rutacearpine suppresses inducible nitric oxide synthase expression and nitric oxide (NO) production by downregulating NF-κB activity in lipopolysaccharide (LPS)-stimulated RAW264.7 macrophages. Treatment with rutacearpine suppressed inducible nitric oxide synthase expression and nitric oxide (NO) production by downregulating NF-κB activity in lipopolysaccharide (LPS)-stimulated RAW264.7 macrophages. Rutacearpine acts by inducing the expression of home oxygenase-1 (HO-1) in a dose- and time-dependent manner. The signaling pathway involved in rutacearpine-mediated HO-1 induction included Ca2+/calmodulin-dependent protein kinase II (CaMkII) and extracellular signal regulated kinase 1/2 (Erk1/2). Furthermore, the CaMkII-Erk1/2 cascade targets the transcription factor, NF-E2-related factor-2 (Nrf2). Taken together, our results demonstrate that rutacearpine-induced expression of HO-1 is mediated by the Ca2+/CaMkII-Nrf2-HO-1 pathway and inhibits LPS-induced inflammation in RAW264.7 macrophages.

Cultivated Ginseng Inhibits Tarc Expression by Suppressing the Activation of NF-κB and STAT1 in Human Keratinocyte Cells.

B. Parki, J. Choi1, Y. Chung2 and H. Jeong1, 1Pharmacy, Chungnam National University, Daejeon, Republic of Korea; 2International University of Korea, into, Republic of Korea.

Overexpression of the HER2 gene causes many cancer types and has been reported to enhance cell proliferation, tumor growth, angiogenesis, and metastasis. This study investigated the molecular mechanisms by which mollugin exerts anti-tumor effect in HER2-overexpressing breast and ovarian cancer cells. Mollugin exhibited potent inhibitory effects on cancer cell proliferation, especially in HER2-overexpressing SK-BR-3 human breast cancer cells and SK-OV-3 human ovarian cancer cells in a dose- and time-dependent manner. Additionally, caspase-3 activity and PARP cleavage were significantly upregulated in HER2-overexpressing SK-BR-3 and SK-OV-3 cells treated with mollugin. Mollugin treatment caused a dose-dependent inhibition of HER2 gene expression at the transcriptional level, potentially in part through suppression of NF-κB activation. Moreover, mollugin inhibits cyclin D1 expression by downregulating HER2 activation and consequently reducing PI3K/Akt signaling. These results suggest that mollugin is a novel modulator of the HER2 pathway in HER2-overexpressing cancer cells with a potential role in the treatment and prevention of human breast and ovarian cancer with HER2 overexpression.

Multiple Signaling Pathways Involved in Suppression of MDR1 by Mollugin from Rubrica cordifolia.

T. Tran, H. Kim, M. Do, S. Jin, E. Shim, H. Han, M. Na and H. Jeong, Pharmacy, Chungnam National University, Daejeon, Republic of Korea.

Multidrug resistance (MDR) is known to be a serious problem in cancer treatment and has been identified as a negative prognostic factor in malignancies. This study investigated mollugin, purified from roots of Rubrica cordifolia L., down-regulated MDR1 expression in MCF-7/adriamycin (MCF-7/adr) cells, human breast multidrug-resistant cancer cell. Mollugin treatment significantly inhibited MDR1 expression by blocking MDR1 gene transcription. Mollugin treatment also significantly increased intracellular accumulation of the fluorescently-tagged P-gp substrate, rhodamine-123. The suppression of MDR1 promoter activity and protein expression was mediated through mollugin-induced activation of AMPK-activated protein kinase (AMPK) and inhibited NF-κB as well as CREB activation.
These results suggest that mollugin treatment enhanced suppression of P-gp expression by inhibiting the NF-κB signaling pathway and attenuating CRE transcriptional activity through AMPK activation.

Liver Kinase B1 Is Required for Phyllin-Induced AMPK Activation in Human HepG2 Hepatocytes.

H. Han, M. Do, H. Kim, T. Khanal, T. Tran, M. Na and H. Jeong, Pharmacy, Chungnam National University, Daejeon, Republic of Korea.

Phyllin, an active constituent found in certain functional foods, has anti-obesity activity in vivo. This study investigated the ability of phyllin to induce AMP-activated protein kinase (AMPK) in human HepG2 hepatocytes. Phyllin strongly activated the phosphorylation of AMPKα at Thr172 in HepG2 cells under normal glucose condition. Additionally, the phosphorylation of AMPKε at Thr172 and ACC at Ser79 was significantly suppressed in cells treated with high glucose, phyllin dose-dependently recovered the phosphorylation of AMPKα at Thr172 and the downstream target acetyl-CoA carboxylase (ACC) phosphorylation at Ser79 in HepG2 cells pretreated by phyllin. Moreover, phyllin significantly stimulated the phosphorylation of liver kinase B1 (LKB1) at Ser428 in a time-dependent manner, with a time course matching that of AMPKε phosphorylation at Thr172. In addition, the defect of phyllin-stimulated AMPKε activation in HeLa cells deficient in liver kinase B1 (LKB1) or in siRNA LKB1-treated HepG2 cells, suggesting that LKB1 is required for phyllin-induced AMPK activation. These results indicate that anti-obesity effects are mediated, at least in part, by the activation of LKB1/AMPK.

Prevention of Free Fatty Acid-Induced Hepatic Steatosis by S-Allyl Cysteine through AMPK Pathways.

H. Jeong1, Y. Hwang2, H. Kim1, J. Choi1 and Y. Chung1, Pharmacy, Chungnam National University, Daejeon, Republic of Korea; 2Korea International University, Jinju, Republic of Korea.

S-allylcysteine (SAC) is the most abundant organosulfur compound in aged garlic extract (AGE), which has been used to standardize commercial aged extracts. SAC has been reported to have antioxidant, anti-cancer, anti-hepatotoxic and neurotrophic properties. In this study, we provide evidence that SAC prevented free fatty acid (FFA)-induced lipid accumulation and lipotoxicity in hepatocytes. SAC significantly reduced FFA-induced generation of reactive oxygen species, caspase activation, and subsequent cell death. Also, SAC mitigated total cellular lipid and TG accumulation in steatotic HepG2 cells. SAC significantly increased the phosphorylation of AMP-activated protein kinase (AMPK) and acetyl-CoA carboxylase (ACC) in HepG2 cells. Additionally, SAC down-regulated the levels of sterol regulatory element binding protein-1 (SREBP-1) and its target genes, including ACC and fatty acid synthase (FAS). Use of a specific inhibitor showed that SAC activated AMPK via calcium/calmodulin-dependent kinase (CaMKK) and SIRT1. Our results demonstrate that SAC activates AMPK through CaMKK and inhibits SREBP-1-mediated hepatic lipogenesis. The results indicate that SAC inhibit lipogenesis in cultured human hepatocytes, and further suggest that SAC impair triglyceride synthesis in part due to decreased de novo fatty acid synthesis resulting from inhibition on AMPK-dependent FAS and SREBP-1c.

Cyto-/Genotoxic Effect of Extract from Bufo bufo garragasini Cantor in Human T Cell Leukemia Cells.


Cinobufacini (Huachansu), an extract from the skins of Bufo bufo garragasini Cantor and has been widely used as an oriental medicine for cancer therapy. Although numerous experimental studies on cinobufacini have been investigated its anti-cancer properties, there is no information whether cinobufacini has anti-cancer effects on human leukemia. The genotoxicity of cinobufacini has not been determined. The aim of the present study was to examine the cyto-/genotoxic effect of cinobufacini in human T cell leukemia (Jurkat T-cells) and human normal lymphocytes. Jurkat T-cells and normal lymphocytes were treated for 24 or 48 hours with various concentrations of cinobufacini. Cytotoxicity was evaluated by Trypan blue exclusion assays. Apoptotic cell death was detected by propidium iodide (PI) staining, and intracellular reactive oxygen species (ROS) was determined by fluorescent dye 5-(and 6)-141-carboxy-2, 7-dichlorodihydrofluorescein diacetate (DCF-DA; Molecular Probes) using flow cytometry analysis. The genotoxicity of cinobufacini in Jurkat-T cells was measured by comet assay and cytokinesis-block micronucleus assay. Cinobufacini significantly inhibited cell viability in Jurkat-T cells but not in normal lymphocytes in dose- and time-dependent manner. Jurkat T-cells treated with cinobufacini induced significantly higher levels of sub-diploid cells compared with control. The DNA content in the sub-G1 region increased from 0.53 % to 22.2 % and from 0.63 % to 42.25 %, when Jurkat-T cells treated with cinobufacini for 24 or 48 hours, respectively. Cinobufacini treatment resulted in significantly higher levels of intracellular ROS. Cinobufacini also induced significant DNA damage determined by comet assay and cytokinesis-block micronucleus assay. In conclusion, these findings suggest that an extract from Bufo bufo garragasini Cantor may be effective for treatment of leukemia.


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The World Health Organization recognizes the incorporation of plant, animal and/or mineral based products in traditional medicines. Millions of South Africans rely on traditional medicines for their primary health care needs. Heavy metal toxicity related to the use of traditional medicine has been reported worldwide. Although minerals are widely available at South African traditional medicine markets, there is insufficient pragmatic information regarding them. Thus the aim of this study was to identify the commonly used crude metal and crystalline salts in the South African traditional medicine trade. Premarket surveys indicated commonly available minerals. These minerals were then purchased from a rural traditional medicine market. Information regarding the colloquial name, mode of administration as well as the price and weight were recorded. Scanning electron microscopy-energy dispersive X-ray was used to determine the composition of the unknown medicinally used powdered products. The elemental spectra was viewed using a Leo 1450 SEM. Mineral salts identified included ammonium chloride, calcium sulphate and sulphur powder. If not handled correctly, such salts can cause numerous health issues including eye irritation and dermal toxicity. Among the available metal salts were potassium dichromate, a Class 1 carcinogen, which is routinely taken both orally and as an enema, and potassium permanganate. Also very concerning was the finding of liquid mercury documented as being ingested for female reproductive complaints. The study is the first to classify and document the commonly available minerals used in South African traditional medicine. Identifying such products has revealed actual and present hazards in the South African traditional medicine trade. Training programmes are underway to alert traditional healers to the risks involved in handling and consuming such potentially harmful substances as well as to provide healthcare workers with algorithms for metal poisonings.

Safety Assessment of Botanical Extracts in Cosmetics.

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It’s a worldwide trend that customers prefer to use personal care products (PCP) containing natural and organic ingredients from botanicals. However natural does not equal safe. In fact, various preparations and complex compositions of plants have great potential to cause adverse effects such as irritation, sensitization and systemic toxicity. The current dilemma is that there is no official guideline to evaluate botanicals in PCP and the chemical assessment cannot be easily adapted to the safety assessment of botanical extractive substances. In line with the guideline issued by Scientific Committee on Consumer Safety in EU, we proposed a strategy to guide the safety evaluation of botanical extracts. The first and foremost is a good characterization of the botanicals which include species, geographic origin, growing and harvesting conditions, manufacturing process, profile of macro and micrometals, analytical markers, chromatographic fingerprint, known toxins, etc. Other commonly checked points are UV absorptive capability, 26 major allergens, residual solvents, and/or metal based products in traditional medicines. Millions of South Africans rely on traditional medicines for their primary health care needs. Heavy metal toxicity related to the use of traditional medicine has been reported worldwide. Although minerals are widely available at South African traditional medicine markets, there is insufficient pragmatic information regarding them. Thus the aim of this study was to identify the commonly used crude metal and crystalline salts in the South African traditional medicine trade. Premarket surveys indicated commonly available minerals. These minerals were then purchased from a rural traditional medicine market. Information regarding the colloquial name, mode of administration as well as the price and weight were recorded. Scanning electron microscopy-energy dispersive X-ray was used to determine the composition of the unknown medicinally used powdered products. The elemental spectra was viewed using a Leo 1450 SEM. Mineral salts identified included ammonium chloride, calcium sulphate and sulphur powder. If not handled correctly, such salts can cause numerous health issues including eye irritation and dermal toxicity. Among the available metal salts were potassium dichromate, a Class 1 carcinogen, which is routinely taken both orally and as an enema, and potassium permanganate. Also very concerning was the finding of liquid mercury documented as being ingested for female reproductive complaints. The study is the first to classify and document the commonly available minerals used in South African traditional medicine. Identifying such products has revealed actual and present hazards in the South African traditional medicine trade. Training programmes are underway to alert traditional healers to the risks involved in handling and consuming such potentially harmful substances as well as to provide healthcare workers with algorithms for metal poisonings.
1797 Effects of Crocin and Safranal, Constituents of Saffron, in 22Rv1 Prostate Cancer Cells.

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Saffron extracts have induced apoptosis, cell cycle arrest, inhibited cellular proliferation, and tumor progression in various cancer cell lines. We are interested in studying the potential chemopreventative effects of saffron especially as it relates to prostate cancer. Recognized active constituents of saffron are crocin and safranal. Cytoxicity of safranal was investigated using the androgen responsive 22Rv1 prostate cancer cell line. The cytotoxicity IC50 of safranal at 24 hr was 141 μM using the tetrazolium dye assay (XTT). The assay was incompatible with crocin. Using the Caspase-Glo® 3/7 assay system, it appears that apoptotic mechanisms were involved in safranal’s cytotoxicity because after 6 hr of exposure, the EC50 of apoptosis was similar to the cytotoxicity IC50. Safranal’s antioxidant activity as measured by a 2,7’-dichlorodihydrofluorescein diacetate assay indicated decreased reactive oxygen species formation. Ongoing studies are investigating the potential of saffron to also inhibit prostate cell invasion and migration in vitro. (Supported by Saudi Arabian Ministry of Higher Education and Salman bin Abdulaziz University)

1798 Kahweol Induces Apoptosis Through Inhibition of STAT3 Phosphorylation in Human Lung Adenocarcinoma A549 Cells.

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Epidemiological studies have shown that unfiltered coffee consumption is associated with a low incidence of cancer. Kahweol, the coffee-specific diterpene, has been reported to have anti-carcinogenic properties. Animal studies have shown that kahweol has a chemopreventive effect of coffee. However, the precise underlying protective mechanisms are poorly understood. In this study, the apoptotic effect of kahweol in human lung adenocarcinoma A549 cells was investigated. In cell viability assays and cell proliferation assays, kahweol exhibited anti-proliferative and pro-apoptotic effects on A549 cells in a time- and dose-dependent manner. Kahweol considerably inhibited the expression of Bel-2 but increased that of Bax; it also stimulated the cleavage of caspase-3 and poly ADP-ribose polymerase. In addition, kahweol inhibited the expression of Bcl-2 but increased that of Bax; it also stimulated the cleavage of caspase-3 and poly ADP-ribose polymerase. The results of the study suggest that kahweol may be a potential chemopreventive agent for lung cancer.

1799 In Vitro Antioxidant Activities of Fractions of Clerodendrum vilicacum Leaf Extract.

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Many diseases are mediated by reactive oxygen species. Clerodendrum vilicacum is a medicinal plant used indigenously in Nigeria for the treatment of some of such diseases. In this study, the antioxidant activities of the hexane, ethyl acetate and methanolic fractions of Clerodendrum vilicacum leaf extract were evaluated in vitro using 2,2-diphenyl-1-picrylhydrazyl-hydrate (DPPH), superoxide anion and hydrogen peroxide scavenging assays. The antioxidant components of the extract fractions were also determined. The results showed that the methanolic fraction had the highest concentrations of vitamin C, vitamin E, selenium, phenols and flavonoids. Moreover, the methanolic fraction of the extract had the highest free radical scavenging activities against DPPH, superoxide anion and hydrogen peroxide with EC50 values of 0.65 ± 0.01, 0.60 ± 0.01 and 0.60 ± 0.01 mg/ml respectively. It also had the highest reducing power against ferric ion. The results of the study suggest that the methanolic fraction of Clerodendrum vilicacum leaf extract may be a potent source of antioxidant compounds which may be useful for the prevention and treatment of reactive oxygen species-mediated diseases.

1800 Comparative Pharmacodynamic Effects of Rituximab-EU (MabThera®) and Rituximab-Pfizer in Cynomolgus Monkeys.

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Background: Rituximab-Pfizer is in Phase 1/2 in RA patients as a potential biosimilar. Analytical and functional characterization has demonstrated in vitro similarity to the licensed rituximab products. Methods: The pharmacodynamic effects of rituximab-Pfizer and rituximab-EU were compared in sexually-mature cynomolgus monkeys in single-dose PK and repeat-dose toxicity studies; both had a 13-week postdose observation period to assess B cell repletion. Peripheral blood lymphocytes were evaluated by flow cytometry, and spleen and axillary and mesenteric lymph nodes (repeat-dose only) were examined microscopically by CD20+ immunohistochemistry. Results: B cell effects were similar in magnitude and time course between rituximab-Pfizer and rituximab-EU. Marked to complete depletion of peripheral blood B cells occurred on Day 4 (the first time point evaluated) in both studies. In the single-dose study, repletion of B cells began on Day 15 (2 mg/kg) and Day 29 (10 and 20 mg/kg) and continued through Day 92, with a subset of animals in each group near or above the pre-dose values at the end of the 13-week observation period. In the repeat-dose study, peripheral B cell depletion persisted through Day 30 of the dosing phase and was associated with lower splenic weights, decreased lymphoid follicle cellularity and decreased CD20+ cells in lymphoid tissues. After repeat-dose, partial repletion of peripheral blood B cells was noted in recovery animals by Day 92 of the recovery phase. Complete histopathological recovery occurred in 3 recovery animals (2 rituximab-Pfizer and 1 rituximab-EU). Lymphoid cellularity and CD20+ cells were increased in the remaining 9 recovery animals (relative to the dosing phase results), indicating partial recovery. Conclusion: The magnitude and time course of B cell depletion and repletion were similar between rituximab-Pfizer and rituximab-EU and were consistent with the expected pharmacology of anti-CD20 monoclonal antibodies and the reported innovator data.

1801 Brain Microhemorrhage Assessment of an Antiamyloid Beta Peptide (Aβ) Monoclonal Antibody (mAb) Using a Transgenic Mouse Model of Alzheimer’s Disease (AD).

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Solanezumab (SLZ), a humanized mAb against Aβ peptide, is being developed for the treatment of AD. SLZ recognizes a mid domain epitope of the Aβ peptide with high affinity and selectivity to soluble monomer. The potential to cause cerebral amyloid angiopathy (CAA)-associated microhemorrhage (MH) was studied in an aged transgenic mouse (APP/V717F) model of AD using a murine surrogate of SLZ, m266.2. Critical study design factors included the use of: 1) age-optimized mice (>21 months) known to have established cerebral amyloid prior to treatment; and thus potentially susceptible to CAA-MH; 2) sufficient numbers (>21 months) known to have established cerebral amyloid prior to treatment; and thus potentially susceptible to CAA-MH; 3) a dose projected to achieve a near-maximal pharmacologic response; 4) a positive control mAb against an N-terminal epitope of Aβ peptide (3D6); 5) meticulous attention to brain collection and processing; 6) expanded brain sectioning and specialized microscopy; and 7) examinations by primary and peer-review pathologists experienced in neuropathologic assessments. Groups of mice received either vehicle control, 50 mg/kg m266.2, or 50 mg/kg 3D6 by weekly ip injection for 4 months (curtailed from the original 6-month duration due to age-related mortality). Plasma exposure to m266.2 and 3D6 was demonstrated at the end of the study. Multiple histologic brain sections stained with Perls’ Prussian Blue/DAB-enhanced Perls from each animal were evaluated to score siderophages resulting from hemorrhage, and multiple H&E stained slides were examined with both brightfield and epifluorescent illumination for the presence of other changes in the brain. While the positive control 3D6 elicited the expected robust microhemorrhage response associated with vascular degeneration, the brains of mice given
1802 Safety Assessment of REGN1001 a Monoclonal Antibody Against Angiopoietin-Like 4 (AngPTL4) in Cynomolgus Monkey Toxicology Studies Under Normal and High-Fat Diet Feeding Regimens.

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REGN1001 is an antibody directed against Angiopoietin-like 4 (AngPTL4) and blocks its ability to inhibit lipoprotein lipase (LPL), an enzyme that hydrolyzes triglycerides (TG) and hence leads to a reduction in plasma TG levels. Evidence for the role of AngPTL4 as a regulator of TG metabolism has been obtained in AngPTL4 gene knockout studies in which profound decreases in TG levels were observed. Several studies have also indicated that AngPTL4 ablation in rodents either via AngPTL4 gene KO or Ab inhibition during high-fat diet (HFD feeding) regimens can lead to the prominent appearance of foamy macrophages and severe inflammatory changes in mesenteric lymph nodes (MLN). To better predict the safety profile of REGN1001 in hyperlipidemic patients (prone to ingest a high fat diet), a toxicology program evaluating REGN1001 intravenous dosing under normal diet and high-fat diet conditions was undertaken in 2 separate 13-week cynomolgus monkey toxicology studies. The first study utilized a normal (high-fat) feeding regimen; the second study used a HFD regimen that enables a western diet. Consistent with the data described in rodents, only the HFD regimen resulted in foamy macrophages in the MLN in monkeys. After a 15-week treatment free recovery period, during which half the animals were withdrawn from HFD, the incidence/severity of foamy macrophages in MLN was decreased relative to those maintained on the HFD regimen. These studies for the first time demonstrate in primates that AngPTL4 inhibition under HFD fed conditions can result in the presence of foamy macrophages in the MLN. In addition, these toxicology studies demonstrate the importance of incorporating human disease factors into animal models for safety testing. Given these data, the likelihood of a similar, clinically-relevant toxicity occurring in hyperlipidemic humans receiving REGN1001 or other ANGPTL4 inhibitors cannot be discounted.

1803 13-Week Toxicity Study of an Anti-Interleukin-6 Antibody (MEDI5117) by Every Other Week Intravenous or Subcutaneous Injection to Cynomolgus Monkeys.

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MEDI5117 is a human immunoglobulin G1 (IgG1)-YTE (triple mutation) monoclonal antibody that binds to interleukin 6 (IL-6) with sub-pM affinity and neutralizes it by preventing binding to the interleukin 6 receptor (IL-6R). The Fc domain of MEDI5117 contains the YTE mutation that increases MEDI5117 Fc-binding to the neonatal Fc receptor (FcRn) at pH 6.0 and thus increases the antibody half-life. To support the first in human clinical study, the nonclinical safety of MEDI5117 was assessed in a GLP repeat IV or SC administration study in cynomolgus monkeys when given by IV bolus injection (15 or 100 mg/kg) or SC injection (15 or 50 mg/kg) every other week for 13 consecutive weeks (7 total dose administrations). Administration of MEDI5117 by SC injection did not result in adverse findings. Administration of MEDI5117 by IV bolus injection resulted in the unscheduled euthanasia of one 15 mg/kg animal that had apparent and treatable anaphylactic reactions to the two previous MEDI5117 dose administrations. There were no similar findings for the other 15 mg/kg IV animals or for animals administered the higher 100 mg/kg IV dose level, and the data demonstrated decreased observed pharmacodynamic and toxicokinetic profiles from this animal as compared to other 15 or 50 mg/kg IV animals receiving anti-drug antibodies. Since this was most likely related to the formation of anti-drug antibody that has been recognized by regulatory agencies as not predictive of immunogenicity in humans and there was a lack of any other findings which is consistent with the toxicology profiles reported for other anti-IL-6 antibodies (Martin et al., 2004 and Femta Pharmaceuticals IND Submission Press Release), this adverse event was ruled as not-related into the NOAEL determination. Therefore, the NOAEL from the GLP study was 100 mg/kg for animals dosed IV and 50 mg/kg for animals dosed SC, the highest doses tested.


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Rheumatoid arthritis is a progressive and disabling autoimmune disease characterized by inflammation of the joints, with subsequent long term structural damage, chronic pain, and limited daily activity. Despite the availability of a variety of effective therapies, a significant portion of patients fail to achieve optimal outcomes including clinical remission and ongoing low disease activity. MedImmune is currently pursuing development of mavrilimumab (CAM-3001), a human monoclonal antibody (mAb) targeting GM-CSF receptor alpha, as a novel treatment for RA. GM-CSF plays a central role in the pathogenesis of rheumatoid arthritis (RA) through the activation, differentiation, and survival of macrophages and neutrophils. The nonclinical safety of mavrilimumab was evaluated in several studies in cynomolgus monkeys as the pharmacologically relevant species. Overall, the nonclinical safety results supported the continued clinical development of mavrilimumab. In clinical studies in RA patients, mavrilimumab has demonstrated good clinical activity with adequate safety to support further clinical development. A Phase 2b study of mavrilimumab in subjects with RA is in progress.

1805 Preclinical Safety and Pharmacodynamics of an Anti-GM-CSF Monoclonal Antibody in Cynomolgus Macaques.


Administration of anti-GM-CSF monoclonal antibodies (mAb) have been proposed as potential therapeutics for inflammatory diseases. In addition to its role in immunity and inflammation, GM-CSF is required for proper maintenance of surfactant catabolism by pulmonary alveolar macrophages (PAM). Humans, loss of GM-CSF signalling results in deregulated surfactant turnover and can manifest as pulmonary alveolar proteinosis (PAP). Therefore, specialized assessments were included in a toxicology study to assess the potential for an anti-GM-CSF mAb to affect surfactant catabolism. The mAb was given intravenously at 0.25 or 50 mg/kg once weekly for 4 weeks or subcutaneously at 0.25, 5 or 50 mg/kg for 13 weeks. In addition to standard toxicity endpoints, bronchoalveolar lavage fluid (BALF) was assessed for cell differentials, intracellular lipids and surfactant protein-D (SP-D) levels. Serum SP-D levels were also evaluated. The mAb was well tolerated with no clinical signs of toxicity. There were no adverse effects on clinical or anatomic pathology. Non-adverse effects in the lung included a dose related increase in the percentage of enlarged PAM, increased numbers of lipid containing PAM and increased BALF SP-D levels. These minor changes may be related to changes in surfactant catabolism but were not accompanied by clinical or histopathological findings indicative of PAP. Serum SP-D values were comparable across groups, consistent with absence of PAP as determined by histopathology. In ex vivo assays, GM-CSF-induced CD11b, pStat5 and proliferation of TF-1 cells were inhibited in a dose dependent manner consistent with expected pharmacodynamic properties. Data for GM-CSF induced CD11b, pStat5 and proliferation were conducted at concentrations of 0.25 mg/kg, which was significantly decreased by the formation of anti-drug antibodies which correlated with decreased activity in the ex vivo assays. In conclusion, anti-GM-CSF mAb was well tolerated by cynomolgus monkeys and produced expected pharmacological effects that were reversible upon treatment cessation.

1806 Nonclinical Safety Evaluation of Anti-PD-L1 (MDP3280A) in Mice and Cynomolgus Monkeys.


Programmed cell death 1 (PD-1) is a receptor expressed on T cells following activation and binding to its ligand, PD-L1, down-regulates the quality and magnitude of T-cell responses. Many neoplastic cells express PD-L1 and evade destruction by the immune system. MDP3280A, an effectors (FcγR-binding deficient) human IgG1 mAb that blocks PD-1/PD-L1 interactions, is in development as a potential therapy for solid tumors. The nonclinical safety program included an in vitro cytokine release assay with human PBMCs, a tissue cross reactivity study, an exploratory 15-day repeat-dose study in C57BL/6 and CD-1 mice, and an 8 week repeat dose safety study in cynomolgus monkeys. MDP3280A did not induce cytokine release from human PBMCs. MDP3280A-specific staining was detected in the lymph node of monkey tissues and the placenta, lymph node, tonsil, and thymus of
human tissues. In vivo, no adverse drug related changes in immunologic endpoints were observed in either species. Drug-related findings were consistent with the expected pharmacology following PD-1/PD-L1 pathway inhibition. In mice, this included reversible increases in splenic weights in both strains, attributed to an enhanced immune response to a heterologous mAb. Minimal sciatric neuropathy with inflammation was observed in C57BL/6 mice only, a strain expressing the MHC H-2b haplotype that in PD-1-deficient mice, develops spontaneous autoimmune peripheral neuropathy. In monkeys, arthritis/periarteritis in parenchymal and/or tubular organs was observed, which is a recognized spontaneous inflammatory condition in this species, and may reflect an MPDL3280A-related enhancement of a pre-existing condition. A high incidence of anti-therapeutic antibodies (50/56 (89%)), which had no consistent impact on exposure, and minimal SC injection site reactions were also attributed to MPDL3280A-enhanced immune responses. These findings are consistent with PD-1/PD-L1 inhibition and identify heightened immune responses and the potential to increase autoimmune liabilities in predisposed individuals as possible safety risks in patients.

**1807 Investigation of the Potential Role of Immunogenicity in Abatacept-Related Lymphocytic Inflammation in Adult and Juvenile Rats.**

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Abatacept is a fusion protein of a human IgG1 Fc and the extracellular domain of the human CTLA-4 that inhibits T-cell activation. Species specific lymphocytic inflammation of the thyroid and pancreatic islets was observed in adult and juvenile rats treated with abatacept subcutaneously (SC) every 3 days for 3 months at pharmacologic doses of 65 and 20 mg/kg/day respectively. Studies in adult and juvenile rats were performed to assess a potential relationship of immunogenicity to lymphocytic inflammation observed in juvenile and adult rats. Abatacept was administered SC at 0 (control) or 0.03 mg/kg (a subpharmacologic dose) to adult rats (20/sex/group) on Days 1 and 29, or to juvenile rats (20/sex/group) on Days 1 (postnatal day 28), 15, 29, and 43. Anti-drug antibodies (ADA) were measured on Days 1 and 29, or to juvenile rats (20/sex/group) on Days 1

**1808 Assessment of Bio comparability of NU100 and Betaferon in Cynomolgus Monkeys.**

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Betaferon is a marketed recombinant human interferon beta-1b (IFN beta-1b) for treatment of relapsing-remitting multiple sclerosis (RRMS). NU100 is an improved recombinant human IFN beta-1b produced using proprietary manufacturing technology. NU100 is aggregate-free and HSA-free in its formulation. In a GLP monkey study for NU100 safety assessment, male and female cynomolgus monkeys were assigned to 2 groups (4 animals/sex/group), and received once every other day NU100 or Betaferon at a dose level of 0.06 mg/kg/dose for 15 days. Animals were monitored for safety; blood samples were collected to determine serum levels of IFN beta-1b, neopterin (a biomarker for IFN-beta-1b pharmacodynamic profile), and anti drug antibodies (ADA). A NU-100- or Betaferon-related adverse safety signs occurred within these 15 days. IFN-beta-1b levels peaked approximately 3 hours postdose; Cmax and AUCCall for NU100 and Betaferon were 23.7 (CV 49.1%) and 14.3 (CV 48.0%) ng/mL and 225 (CV 51.6%) and 157 (CV 46.6%) h•ng/mL at steady state, respectively, and were not statistically significantly different. Neopterin levels peaked 24 hours postdose on Day 1; Emax and AUCECall for NU100 and Betaferon were 5.4 (CV 41.4%) and 6.29 (CV 31.7%) nMol/L and 65.9 (CV 45.2%) and 75.7 (CV 39.5%) h•nMol/L, respectively; whereas Neopterin concentrations reached steady state on Day 15; Emax and AUCECall for NU100 and Betaferon were 1.71 (CV 78.1%) and 1.03 (CV 107.2%) nMol/L and 21 (CV 77.3%) and 35.6 (CV 7.9%) h•nMol/L, respectively, and were not statistically significantly different. This reduced response on Day 15 to treatment was likely due to the production of anti-drug antibody (ADA), with 5 of 8 animals in the NU100 group positive for ADA. In conclusion, NU100 and Betaferon had comparable safety, toxicokinetic, and pharmacodynamic profiles in cynomolgus monkeys.

**1809 Preclinical Development and Safety Assessment of the First Inhaled Nanobody ALX-0171.**

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Nanobodies are therapeutic proteins based on the smallest functional fragments of naturally occurring heavy chain only antibodies. The trivalent Nanobody ALX-0171 targets the respiratory syncytial virus (RSV) with high specificity and potency. It has the potential to be effective in prevention and treatment of RSV infection, a cause of severe upper and lower airway inflammation in susceptible populations. The medical need is high in young children, with 0.3 million patients younger than 5 years hospitalised every year. ALX-0171 is formulated as a nebulizer solution for pulmonary inhalation as clinical route of administration. Using a vibrating mesh nebuliser, a droplet size ≤3.4 μm (MMAD) was achieved, while the drug's stability was confirmed. The preclinical package consisted of 2-week repeated dose toxicity studies via iv administration or inhalation in adult rats, a respiratory safety pharmacology study in rats via single inhaled dose and a cardiovascular safety pharmacology study in dogs (iv, single ascending dose). No drug- or immunogenicity-related safety findings were observed. A standard BCOP assay demonstrated the non-corrosive nature of the compound. In addition, a cotton rat disease model was performed to assess safety, tolerability and efficacy of ALX-0171 (intratracheal delivered). A dose-dependent reduction of viral transcripts following viral re-challenge in lung tissue of RSV-infected cotton rats was demonstrated as efficacy marker. A significant improvement of infection-related events (body weight stagnation, organ weight, BALT inflammatory cell counts) was demonstrated without signs of immune-induced events upon ALX-0171 administration. Treatment-induced anti-drug antibodies were measured in BALF and plasma and results indicated a mild immunogenicity response. It can be concluded that ALX-0171 was well-tolerated in safety assessments in preclinical species. ALX-0171 successfully completed a phase I clinical trial.

**1810 The Occurrence of Microscopic Vacuoles in Toxicology Studies with Marketed PEGylated Proteins Is Associated with High Doses, High Clinical Multiples, and Accumulation.**

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To obtain a more complete understanding of the clinical significance of microscopic vacuoles observed in toxicology studies with marketed pegylated proteins, nonclinical programs were reviewed from FOI data (publicly available BLA reviews) for Omontys, Krystexxa, Cimzia, Micrera, Macugen, Somavert, Pegafus, Neulasta, PegIntron, and Adagen. Cumulative toxicity doses for conjugate, PEG, and protein were calculated for each study and compared with cumulative recommended clinical doses over the same interval. None of the studies included PEG control groups. Microscopic pathology evaluations across programs were part of GLP toxicity studies and were performed on formalin/immersion-fixed tissues. There was no indication of whole-body perfusion techniques. Microscopic vacuoles were noted in toxicity studies with Omontys, Krystexxa, Cimzia, Macugen (IV but not intravitreal), Somavert, and Neulasta but not for Micrera, Micrera (intravenous), Pegafus, PegIntron or PEG1KD. Across programs, their appearance and reversibility were dose-related and associated with large cumulative PEG/conjugate doses, short inter-dose intervals, longer study durations, drug accumulation, and large clinical multiples. At high cumulative doses (up to 14,000-fold the recommended clinical dose based on mg/m² and PEG doses of 2720 mg/m²), vacuoles were noted in the macrophage phagocytosis system and, in some instances, choroid plexus, uterus, ovary, pituitary, and adrenal cortex. Nearly all of the high doses associated with vacuoles exceeded high-dose guidelines per ICH S6 Addendum. In conclusion, the appearance of microscopic vacuoles in toxicity studies with marketed pegylated proteins appears related to high cumulative doses.
1811 TAS-116, an Oral Highly Potent HSP90α/β Selective Inhibitor, Leads Minimized Ocular Toxicity in Both Albino and Pigmented Rats.

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Sponsor: M. Takahashi

BACKGROUND: Heat Shock Protein 90 (HSP90) is a key chaperon which has a best-in class HSP90 inhibitor with minimized ocular toxicity.

RESULTS: When TAS-116 administered orally for 14 days, no dose-related change was revealed in ophthalmological examination in albino rat. In histopathology, reference HSP90 inhibitors (AUY922 and 17-DMAG) caused degeneration and/or disarrangement of photoreceptor cells and increase in TUNEL positive apoptotic cells in retinal outer nuclear layer (ONL). On the other hand, TAS-116 demonstrated no histological changes or increase in TUNEL positive cells in ONL in albino rat. When all compound administered intravenously in albino rat, AUY922 and 17-DMAG showed greater exposure in retina compared to plasma, whereas TAS-116 showed less distribution in retina than in plasma. In addition, oral administration of TAS-116 demonstrated less retinal distribution and did not accumulate in retina after 2 week repeated dosing. In contrast, TAS-116 indicated a much higher distribution in subcutaneously implanted tumor over retina in rat model. Furthermore, TAS-116 had no melanin affinity because TAS-116 did not induce the retinal toxicity and also showed less distribution in retina in pigmented rat.

CONCLUSION: TAS-116 does not induce ocular toxicity in both albino and pigmented rat. This is probably due to less distribution in retinal tissue of TAS-116. These unique profiles of TAS-116 indicate that TAS-116 has a potential to be a best-in class HSP90 inhibitor with minimized ocular toxicity.

1814 Preclinical Safety Evaluation of JNJ-35815208, a Selective Estrogen-Related Receptor Alpha Modulator.


Sponsor: M. Takahashi

Estrogen-related receptor-α (ERRα) is an oral nuclear receptor that has emerged as a novel therapeutic target for the treatment of type II diabetes and cancer. Here we describe non-clinical safety evaluation in rats and dogs of a novel and selective ligand, JNJ-35815208, for ERRα as a potential anti-diabetic agent. Following single oral administration in rats, mortality was observed at ≥ 1000 mg/kg. Repeated dose for 14 days at doses 80 and 200 mg/kg/day was well tolerated at doses up to 40 mg/kg. At 200 mg/kg/day, microscopic findings were observed in the testes and epididymides, characterized by mild bilateral degeneration/atrophy of the seminiferous tubules, minimal bilateral degeneration of the germ cells, luminal cellular debris, and oligospermia in the epididymides. In the 4-week rat study at 1, 6, or 40 mg/kg/day, there were no toxicological findings. In dogs, following single-dose administration at 50 and 250 mg/kg, as well as during the 5-day repeat dose at 150 and 250 mg/kg/day, the primary finding was emesis noted at all dose levels. In the four-week dog study at 2, 16 and 75.5/100 mg/kg/day, microscopic findings were noted in the female reproductive organs and the mammary glands at all doses. That resulted in reductions in the size of the ovaries, uterus, and vagina as well as atrophy of the mammary glands. These effects were associated with a persistent anestrus of the reproductive cycle. In male dogs, multifocal atrophy of the glandular epithelium of the prostate gland was observed at the mid- and high doses and small testes accompanied by mild degeneration of the seminal vesicles and vacuolated Sertoli cells at the high dose. Erythroid hypopcellularity in the bone marrow was noted in males at 16 mg/kg/day and in both sexes at 75.5/100 mg/kg/day, with corresponding decreases in RBC, hemoglobin, hematocrit and reticulocyte counts at the high dose. Overall, these data suggest ERRα may play a role in both male and female reproductive organs as well as in the bone marrow.
implantation of human liver and thymic tissue underneath the kidney capsule and CD34+ hematopoietic stem cell transplantation. Approximately 12-16 weeks following surgery, the mice have an engrafted functional human immune system, achieve 20-25% humanization in peripheral blood and are suitable for studies. This approach potentially offers the ability to test for efficacy and safety of drug products in an in vivo model of the human immune system. If proven reliable via testing of clinically available biologics, this animal model could be a powerful tool in drug testing.

In order to begin assessing the ability of this model to predict uniquely human immune responses, we tested two forms of interferon-β (IFN-β) currently marketed in the USA using clinically relevant dosing regimens and routes. Humanized mice were initially given IFN-β1a subcutaneously (sc) at doses of 6.0 μg, 1.5 μg or 0.3 μg once weekly or IFN-β1b at doses of 2.5 μg, 12.5 μg or 25 μg three times weekly for four weeks to determine if acute toxicity would result. A follow-up study using doses of 0.3 μg of IFN-β1a or 5 μg of IFN-β1b for eight weeks assessed serum drug levels, immunogenicity and immune responses. The results of these studies demonstrated that (1) standard drug doses used in IFN-β studies in transgenic IFN-β mice can be toxic to mice with a human immune system, as humanized mice can respond to the drug in a clinically relevant manner; (2) presence of the appropriate human receptors makes drug level assessment possible in humanized mice; and (3) humanized mice respond immunologically to IFN-β.

1816 Evaluation of Antisense Oligonucleotides in Human Phbc and Association of Release of IL-6 In Vitro to Constitutional Symptoms.

H. S. Younis, T. Machemer, D. A. Norris and S. P. Henry, Preclinical Development, ISIS Pharmaceuticals, Carlsbad, CA.

Oligonucleotide based therapeutics produce low-grade non-specific proinflammatory responses that manifest as increased cytokine/chemokine production, splenomegaly and/or lymphohistocytic cell infiltrates in multiple tissues in animal models of toxicity. In humans, antisense oligonucleotides are well tolerated and may produce constitutional effects such as flu-like symptoms and injection site erythema that are typically self-limiting. The objective of this research was to determine if in vitro cytokine production from human peripheral blood mononuclear cells (PBMC) may be used to determine the potential for antisense oligonucleotides (ASO) to produce constitutional symptoms in humans. Fresh PBMC (n=50 donors) were cultured with a selected list of ASOs (ISIS 104838, ISIS 113715, ISIS 325568 and ISIS 353512; 0-80μM) that have been evaluated in humans and produced a broad range of constitutional symptoms (none to moderate). By 24 hr of treatment, the culture supernatant was harvested for measurement of cytokine (IL-6 and IL-10) release. A sigmoidal Emax model was used to determine the Emax and EC50 for the IL-6 response of each ASO. Dose-dependent increases in IL-6 were consistently greater (0.3μM EC50) for ISIS 353512 (the ASO that produced the most pronounced constitutional symptoms in humans) than for the other evaluated ASOs (1.3-3μMEC50 range). The IL-6 Emax/EC50 ratio was 10-fold greater for 353512 than the least responding ASO ISIS 104838, and provided the best fit for differentiating ISIS 353512 from the ASOs generating lesser responses. The release of IL-10 was also greater for ISIS 353512 but was more difficult to quantify given the lack of a dose response and relative low magnitude of change. Collectively, the results suggest that human PBMC may be a viable in vitro model to rank order ASOs for their potential to produce constitutional symptoms in humans and to investigate the mechanisms of ASO mediated proinflammation.

1817 Mechanistic Basis of the Species-Specific Supplemental Activation in Cynomolgus Monkeys with Oligonucleotide Treatment.

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Antisense oligonucleotide (ASO)-mediated alternative pathway of complement (APC) activation is a common, class effect in macaque monkeys at high doses. Activation is due to the interaction between ASO and complement Factor H (CFH), the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the APC. Transient reduction of circulating CFH, the inhibitory protein of the AP...
**1820 Application of Canine Kidney Tissue Slices to Detect the Toxicity of Prototypical Nephrotoxic Agents.**

K. Kowalkowski, M. Kląpczyński, D. J. Cugier, E. A. Blomme, W. R. Buck and M. J. Liguori. *Cellular and Molecular Exploratory Toxicology, AbbVie, Abbott Park, IL.*

Precision-cut renal tissue slices retain the multicellular, structural, and functional features of their original organ and offer a more relevant approach to interrogate toxicity compared to traditional cell-based in vitro systems. Here, we sought to evaluate the utility of this system to detect toxicity induced by prototypical kidney tubular toxicants, including cadmium chloride (CdCl2) and cisplatin. Kidneys from male beagle dogs were cut into 300 μm slices, and cultured at 37°C in an O2-rich atmosphere for 24 to 48 h. Initial experiments optimized the culture conditions to maintain viability to at least 48 h. The slices were immediately treated after isolation with multiple doses of CdCl2, cisplatin, or vehicle. Multiple endpoints of toxicity were evaluated including H&E stained sections, intracellular ATP content, LDH release, total RNA integrity, and a novel endpoint for this type of system, kidney injury molecule-1 (Kim-1) mRNA levels via in situ hybridization. There was a significant elevation of LDH leakage at all doses tested for both compounds. The intracellular level of ATP also declined significantly at 100 μM CdCl2, and 100 μM cisplatin, indicative of substantial cellular damage. Mild to marked degeneration and necrosis of renal tubules was evident beginning at 24 h upon microscopic examination. Kim-1, which has been previously demonstrated as a promising in vivo kidney injury biomarker and is not normally expressed in normal tissue, was clearly overexpressed in CdCl2 treated slices, but not vehicle treated slices, especially along the tubular epithelium. In severely necrotic slices, Kim-1 was undetectable, likely due to RNA destabilization as later confirmed using integrity assessments. These data demonstrate that a renal slice in vitro model can be used to detect potent nephrotoxicants and that Kim-1 should be further explored as a novel in vitro biomarker to monitor toxicity in renal slice systems.

**1821 Investigation of Protein Kinase C Inhibitor-Induced Steroid Hormonal Perturbation in Rat.**


Steroid hormones are crucial endogenous mediators synthesized and secreted into the bloodstream by endocrine glands, such as the adrenal cortex and gonads. They mediate a wide variety of physiological functions. Altered steroid hormone status may affect the progression of various types of cancer. For instance, prepubertal ovarectomy in BFU/Mna rats accelerated the growth of spontaneous thymoma, which could be mitigated by intraperitoneal injection of estrogens. In azaserine-treated rats, the rate of spontaneous pancreatic tumors in males is increased, while orchidectomy reduced the incidence of such neoplasms compared to intact animals.

In a 104-week oral carcinogenicity study in rat with a protein kinase C inhibitor, a treatment related increased incidence of pancreatic acinar adenoma in males and thymoma in females were observed at dosages ≥ 100 mg/kg/day. Since changes in steroid hormones are considered to potentially affect the development of pancreatic and thymus neoplasia in rodents, a 2-week oral mechanistic follow-up study was conducted in rats to investigate this hypothesis. For this purpose, a mass spectrometry method previously developed in our lab for the quantification of steroid hormones in plasma, was further applied to analyze 17 steroids in adrenal, testicular and ovarian tissues. Aldosterone was increased in plasma and adrenals in treated males with a concomitant and pronounced decrease of androstenedione and testosterone in plasma, adrenals and testes. In female rats, progesterone, together with cortisol, showed a trend towards increased concentrations in adrenals and ovaries after treatment, whereas ovarian androstenedione and estrogen levels tended to be reduced. This correlated with a circulating luteinizing hormone decrease in both sexes. In conclusion, the reduction in ovarian estrogen levels might be connected to the thymus neoplasias of the previous study. As androgen levels were found to be decreased, a direct connection to the previously reported pancreatic tumors is unlikely.

**1822 Comparison of In Vivo Central Corneal Thickness (CCT) Measurements in Dogs, Rats, and Nonhuman Primates Using a Handheld Ultrasonic Pachymeter or Specular Microscope.**

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The cornea is responsible for 2/3 of light refraction in the eye and thus is a critical tissue for clear vision. Alterations in corneal thickness have a direct effect on visual acuity and can be caused by alterations in the function of the corneal endothelium, aqueous humor composition, drainage or formation rate, or in the glands producing the tear film. There has recently been an increase in the number of topical ocular therapeutic effects as this method is deemed one of the simplest clinically. As the formulations are in direct contact with the cornea, it can be an intended or unintended target tissue. In non-clinical topical ocular studies, serial in vivo measurement of corneal thickness can provide a rapid empirical insight into the health of the cornea during study conduct and whether or not changes occur. It is reversible. There are several ways to obtain central corneal thickness measurements, two of which are ultrasonic pachymetry (UP) that requires contact with the corneal surface and specular microscopy (SM) which is a non-contact procedure. The data presented represent background thickness measurements and precision between the two methods. In most cases, animals required sedation in order to reduce eye movement. CCT measurements by UP and SM were: dogs (n=10) 548.5±33 μm and 583.5±27 μm; rabbits (n=12) 378±27 μm and 394±16 μm; NHPs (n=10) 422±21 μm and 440±19 μm. Direct comparison showed a difference of 6% between the two methods for dogs, 4% for rabbits and 4% for NHP, with the SM values consistently higher. Precision as determined by 2 separate measurement occasions in dogs was 0.6% for SM and 0.1% for UP. In conclusion, both specular microscopy and ultrasonic pachymetry provide reproducible, precise CCT data in dogs, rabbits and non-human primates allowing for repeat measures during the course of a study when corneal changes are of concern. Care should be taken for interpretation if data are acquired using both methods considering the 4-6% lower results using UP.

**1823 In Vivo Evaluation of the Corneal Endothelial Cell Layer Using Specular Microscopy.**

K. Tenneson and M. Vézina. *Ocular and Neuroscience, Charles River, Montréal, QC, Canada.*

The corneal endothelium is a monolayer of specialized cells whose primary physiological function is to maintain the health and transparency of the corneal stroma. In preclinical ocular and non-ocular studies, pharmacological and/or toxicological effects of a test compound may produce changes in the structure and function of the corneal endothelium. Additionally, intraocular surgery (including induced disease models) or implanted devices can compromise the endothelium and cause corneal edema. Specular microscopy (SM) enables a rapid, minimally invasive, direct evaluation of the cornea, and can provide supplemental information on corneal endotheliopathies, alone or in combination with direct slit-lamp examination on preclinical toxicology studies. SM provides three primary endpoints: number of cells/unit area (density), shape of cells (pleomorphism) and size of cells (poly-megathism). When the endothelium is disrupted, there can be an overall shift in the number of cells, proportion of cell shapes (pleomorphism) and sizes (poly-megathism). In order to characterize the endothelial layer in species regularly used on ocular toxicology studies, dogs, rabbits and non-human primates (NHP) were subjected to SM. Overall, cell density was similar across species (rabbits: 2769 cells/mm2; dogs: 2635 cells/mm2; NHPs: 3058 cells/mm2). Normal endothelial cells have a hexagonal shape, and account for the majority of cells (rabbits: 75%; dogs: 65%; NHPs: 71%). Average cell sizes were 364 um2 for rabbit, 384 um2 for dog, and 328 um2 for NHP. In conclusion, SM is a sophisticated tool that can complement the standard endpoints performed on specialized ocular preclinical studies and the data presented provides background data in rabbits, dogs and NHPs to facilitate the recognition of compound-related changes.

**1824 Therapeutic Effect of MG132 on Diabetic Cardiomyopathy Is Associated with the Suppression of Proteasome Activities: Roles of Nrf2 and NF-kB.**

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MG132, a proteasome inhibitor, can up-regulate nuclear factor erythroid 2-related factor 2 (Nrf2)-mediated anti-oxidation and down-regulate nuclear factor-(NF)-κB-mediated inflammation. The present study was to define whether through above two mechanisms MG132 can provide a therapeutic effect on diabetes-induced cardiomyopathy. To this end, transgenic OVE26 type 1 diabetic mice were used. OVE26 mice develop hyperglycemia at 2 – 3 weeks after birth and exhibit albuminuria and cardiac dysfunction at 3 months of age. Therefore starting at 3 months of age MG132 was used. OVE26 mice develop hyperglycemia at 2 – 3 weeks after birth and exhibit albuminuria and cardiac dysfunction at 3 months of age. Therefore starting at 3 months of age OVE26 diabetic mice were treated intraperitoneally with MG132 at 10 μg/kg body-weight daily for 3 months. At 3 and 6 months of age, cardiac function was measured with M-mode echocardiography. At 6 months, cardiac tissues were subjected to pathological and biochemical examination. OVE26 diabetic mice, but not MG132-treated OVE26 diabetic mice, showed significant cardiac dysfunction, including increased left ventricular systolic diameter and wall thickness and a decreased left ventricular ejection fraction with an increase of heart weight/tibia length ratio. Hearts of OVE26 diabetic mice exhibited structural derangement and remodeling (fibrosis and hypertrophy). In OVE26 diabetic mice,
there was also increased cardiac oxidative damage and inflammation. All of these pathogenic changes were reversed by MG132 treatment, which is associated with a significant suppression of cardiac increase in proteasome activity. In addition, MG132 treatment also significantly up-regulated Nrf2 expression and transcription (shown by increased expression of Nrf2 down-stream antioxidant genes) and down-regulated Ik-B expression and NF-κB nuclear accumulation. These results suggest that MG132 provided a therapeutic effect on diabetic cardiomyopathy in OVE26 diabetic mice possibly via the up-regulation of Nrf2-dependent anti-oxidation and the down-regulation of NF-κB-mediated inflammation.

1825 MG132 Prevents the Progression of Diabetes-Induced Pathological Damage to Aorta Is Associated with Its Up-Regulation of Nrf2 and Its Down-Stream Antioxidant Proteins.

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Endothelial damage and dysfunction are manifested in diabetic cardiovascular complications. Nuclear factor-erythroid 2-related factor 2 (Nrf2) is one of the most important cellular defense mechanisms against oxidative stress by up-regulation of several antioxidants, phase II detoxifying enzymes, and other proteins that detoxify xenobiotics and neutralize reactive oxygen and/or nitrogen species. Deletion of Nrf2 gene significantly enhances the susceptibility of cardiomyocytes to high-level glucose-induced reactive oxygen species generation and cell death. The present study was to define whether induced Nrf2 by MG132 can provide a therapeutic effect on diabetes-induced aortic pathogenic changes. To this end, transgenic OVE26 type 1 diabetic mice and age-matched control mice were used. OVE26 mice develop hyperglycemia at 2–3 weeks after birth and exhibit renal and cardiac dysfunction at 3 months of age, suggesting the induction of diabetic complications. Therefore starting at 3 months of age, OVE26 diabetic mice were intraperitoneally treated with MG132 at 10 μg/kg body weight daily for 3 months. At the end of MG132 treatment, aortas from these mice were morphologically and immunohistochemically examined. Significant increases in the wall thickness and structural derangement of aorta were found in OVE26 diabetic mice, which was accompanied by significant increases in aortic oxidative and/or nitrosative damage (4-HNE as lipid peroxidation and 3-NT as protein nitration), inflammation (TNF-α and PAI-1), and remodeling (CTGF and TGF-β1). These pathological changes were not observed in MG132-treated OVE26 diabetic mice, which was also associated with a significant increase of aortic Nrf2 expression and transcription function. These results suggest the therapeutic effect of MG132 on diabetes-induced aortic pathogenic damages and its association with Nrf2 expression and function.

1826 Characterization of miR-208a Responses in Isoproterenol-Induced Cardiac Injury in Sod2+/- and C57BL/6J Mice.

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The present investigation aimed to characterize miR-208a as a putative biomarker for early cardiac injury in isoproterenol (ISO)-induced acute myocardial damage in mice and to investigate potential strain-dependent effects in response to ISO administration. Plasma from age-matched male C57BL/6J and Sod2 +/- mice treated with a single intraperitoneal (IP) administration of ISO was collected for measuring cardiac troponin I (cTnI) and miR-208a at 3, 6, and 24 hours post injection. Administration of ISO led to increases in both cTnI and miR-208a at all time points tested. However, the magnitude of increase and the temporal release profiles of these biomarkers were different between the two strains. In C57BL/6J mice, cTnI and miR-208a tracked with each other in both magnitude and time, the highest values were seen at 3 hours. By contrast, in Sod2+/- mice, the magnitude of miR-208a was much greater than that of cTnI with the highest values of both biomarkers observed at 6 hours. Similar to C57BL/6J mice, the temporal profile of miR-208a followed that of cTnI in Sod2+/- mice. Histopathological examination of hearts treated with ISO revealed myocardial degeneration at ≥ 3 hours in C57BL/6J mice and ≥ 6 hours in Sod2+/- mice which correlated with the highest concentration of the biomarkers in each strain. The higher systemic exposure of ISO in C57BL/6J mice compared to that in Sod2+/- mice may have contributed to the observed earlier response in C57BL/6J mice compared to Sod2+/- mice.

1827 Polybrominated Diphenyl Ethers Exposure and Intrauterine Growth Restriction: A Case-Control Study in Chinese Newborns.

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BACKGROUND: Intrauterine growth restriction (IUGR) is associated with perinatal morbidity and mortality. It has multifactorial etiology. Along with malnutrition and psychosocial, environmental pollutants, including polybrominated diphenyl ethers (PBDEs), have also been considered to be involved in the etiology of this disease.

OBJECTIVES: This case-control study was performed to assess maternal–fetal exposure to PBDEs and investigate whether in utero PBDE exposure is associated with IUGR.

METHODS: A total of 29 newborn-mother pairs residing in Wenzhou were enrolled in this study during December 2010 and February 2011. Maternal blood and umbilical cord blood (UCB) samples were collected and analyzed for PBDEs by the method of Gas chromatography–mass spectrometry (GC-MS). Conditional logistic regression and Spearman correlation were used to analyze the association between PBDEs exposure and IUGR.

RESULTS: All PBDE congeners in serum were detected except for BDE 138, 183, and 190. BDE 209 was the most abundant congener followed by BDE 207, 208, and 66, with the detection frequencies of 50%, 83%, 74%, and 74%, respectively. The concentrations of BDE 66, BDE 209, BDE 183-209 and 19 PBDEs in UCB are significantly higher in newborns with IUGR than those in controls. BDE 183-209 and 19 PBDEs levels in UCB were inversely associated with birth weight and Quetelet’s index (p=0.008, 0.020 respectively). After controlling for potential confounders, dose-response relationships were observed between IUGR and BDE 183-209 and 19 PBDEs levels in UCB.

CONCLUSION: Only one UCB sample from the control group did not detect PBDEs which might indicate that newborns in China were ubiquitously exposed to PBDEs. Significantly higher PBDEs levels were detected in IUGR cases compared with those in controls. In utero PBDEs (especially high-brominated BDE congeners) exposures were associated with IUGR in a dose-dependent manner. Prenatal PBDEs exposure may be a risk factor for IUGR.


A. J. Schecter1, D. Cherry2, L. S. Hyman3, D. Cheng1, N. Imran4, M. Hommel5, K. Kannan6, L. Wang7, S. H. Yun8, N. Thieue9, B. Specteren1, J. Moye5,4 and L. S. Birnbaum1,8,10. 1University of Texas School of Public Health, Dallas, TX; 2University of the Texas, Tyler, TX; 3The University of Texas Southwestern Medical Center, Dallas, TX; 4New York Health Department, Albany, NY; 5South Dakota State University, Brookings, SD; 6NCI, Bethesda, MD; 7NICHD, Bethesda, MD; 8NIH, Bethesda, MD; 9NIEHS, Research Triangle Park, NC; 10NCI, Bethesda, MD.

The NCS is a prospective health study of 100,000 children (birth to 21 years) and their mothers. We report on a nested, formative study involving measurement of lipid soluble persistent PBDE flame retardants. PBDEs have been associated with various health effects. This study involved 20 mother-child pairs; 20 samples each of maternal third trimester blood, maternal blood at birth, cord blood, and breast milk were tested individually for nine PBDE congeners. Total PBDE was the sum of nine PBDE congeners when quantifiable. Some PBDE congeners were detected in maternal third trimester blood, maternal birth serum, maternal cord blood, and breast milk. BDE 183 and 190. BDE 209 was the most abundant congener followed by BDE 207, 208, and 66, with the detection frequencies of 50%, 83%, 74%, and 74%, respectively. The concentrations of BDE 66, BDE 209, BDE 183-209 and 19 PBDEs in UCB are significantly higher in newborns with IUGR than those in controls. BDE 183-209 and 19 PBDEs levels in UCB were inversely associated with birth weight and Quetelet’s index (p=0.008, 0.020 respectively). After controlling for potential confounders, dose-response relationships were observed between IUGR and BDE 183-209 and 19 PBDEs levels in UCB.

CONCLUSION: Only one UCB sample from the control group did not detect PBDEs which might indicate that newborns in China were ubiquitously exposed to PBDEs. Significantly higher PBDEs levels were detected in IUGR cases compared with those in controls. In utero PBDEs (especially high-brominated BDE congeners) exposures were associated with IUGR in a dose-dependent manner. Prenatal PBDEs exposure may be a risk factor for IUGR.
1829 Effect of Body-Weight Loading onto the Articular Cartilage on the Occurrence of Quinolone-Induced Chondrotoxicity in Juvenile Rats.


Quinolone antibacterial agents have been reported to induce chondrotoxicity in juvenile animals, and the mechanism has not yet been clarified. We have reported that gene expression of tumor necrosis factor receptor superfamily, member 12a (Tnfrsf12a, cell death-related gene), prostaglandin-endoperoxide synthase 2 (Ptgs2, inflammatory response-related gene), plasminogen activator, urokinase receptor (Plaur, stress response-related gene), and matrix metalloproteinase 3 (Mmp3, proteolyis-related gene) was involved in the induction of cartilage lesions of the distal femoral articular cartilage in juvenile rats treated orally with ofloxacin (OFLX). In the present study, the effect of body-weight loading onto the articular cartilage on the occurrence of the cartilage lesions was investigated in male juvenile Sprague-Dawley (SD) rats given OFLX orally once at 900 mg/kg. Just after dosing of OFLX, hindlimb unloading was performed for 0 h, 2 h, 4 h, and 8 h by a tail-suspension method. Animals were sacrificed at 8 h post-dose, and then the distal femoral articular cartilage was subjected to a histological examination and an investigation for gene expression of Tnfrsf12a, Ptgs2, Plaur, and Mmp3 by qRT-PCR analysis. As a result, cartilage lesions and up-regulations of these 4 genes that were seen in rats without the tail suspension were not observed in rats with the 8-h tail suspension, and a tendency to decrease in the incidence of the cartilage lesions and up-regulations of these 4 genes that were seen in rats without the tail suspension were not observed in rats with the 8-h tail suspension, and a tendency to decrease in the occurrence of the cartilage lesions was investigated in male juvenile Sprague-Dawley (SD) rats given OFLX orally once at 900 mg/kg. Just after dosing of OFLX, hindlimb unloading was performed for 0 h, 2 h, 4 h, and 8 h by a tail-suspension method. Animals were sacrificed at 8 h post-dose, and then the distal femoral articular cartilage was subjected to a histological examination and an investigation for gene expression of Tnfrsf12a, Ptgs2, Plaur, and Mmp3 by qRT-PCR analysis. As a result, cartilage lesions and up-regulations of these 4 genes that were seen in rats without the tail suspension were not observed in rats with the 8-h tail suspension, and a tendency to decrease in the incidence of the cartilage lesions and gene expression was noted in a tail-suspension time dependent manner. Our results clearly indicate that body-weight loading onto the cartilage is necessary to induce cartilage lesions and gene expression of Tnfrsf12a, Ptgs2, Plaur, and Mmp3 in juvenile rats treated with OFLX.

1830 Effects of Maternal Exposure to Phthalates and Bisphenol A on Neonates.

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RATIONALE: Phthalate and bisphenol A (BPA) metabolites may have anti-androgenic and pro-inflammatory effects. We hypothesized that maternal exposure to phthalates and BPA in pregnancy is associated with decreased anogenital index in infants and shortened gestation.

METHODS: Urinary concentrations of phthalate and BPA metabolites were measured from 72 pregnant women enrolled in a high-risk obstetric clinic, at the last clinic visit prior to delivery, using high-performance liquid chromatography-tandem mass spectrometry. Anogenital index (AGI) was calculated by normalizing anogenital distance to weight. Gestational age was determined by either sonographic dating or date of implantation. Using linear regression models, we estimated the change in gestational age and anogenital index associated with each interquartile range (IQR) increase in phthalate and BPA metabolite concentration.

RESULTS: IQR increases in urinary mono(2-ethyl-5-hydroxyhexyl) phthalate (MEHHP) and total BPA concentrations were associated with 4.2 and 1.1 day decreases in gestation, respectively. When stratifying by gender, these alterations were found only in male infants. Levels of monoethyl phthalate (MEP) were associated with lower AGI in males, but this difference (0.2 mm) is not clinically significant.

CONCLUSIONS: MEHHP and total BPA were associated with small reductions in gestation, and MEP with a small reduction in AGI, with effects only in males. Increased sensitivity of males to the effects of phthalates and BPA may be related to the roles of androgen precursors in both genital differentiation and in the initiation of labor.

Supported by NIH P30ES005022, R21HD058019 and NJ Dept. of Environmental Protection.

1831 Intravenous Dose Range-Finding Toxicity of Allopregnanolone (ALLO) in Neonatal Dogs.

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Treatment with allopregnanolone (ALLO), a GABA-ergic neurosteroid, may be useful treating Niemann-Pick Type-C (NPC) disease. NPC-C is a rare autosomal recessive neurodegenerative disease caused by genetic mutations, resulting in lysosomal accumulation of unesterified cholesterol and glycolipids. Treatment of neonatal NIH NPC mice with a single s.c dose of ALLO resulted in increased life span, onset delay of neurological symptoms, survival of cerebellar Purkinje and granule cell neurons, and reduced cholesterol and ganglioside accumulation. Previous toxicity studies with ALLO suggested that the presence of isoflurane anesthesia may have contributed to a relatively low MTD of <3 mpk bolus intravenous (IV) injection of unconscious neonatal Beagle dogs. The current study determined the tolerability of ALLO administered IV with extended infusion or bolus injections in conscious pups. Indwelling IV catheters were placed under isoflurane anesthesia. Animals were allowed to recover from anesthesia for ALLO/vehicle treatment. Pups (1-2 per dose, age: postnatal day (PND) 17-PND59) received vehicle 20% hydroxypropyl [β-cyclodextrin (HPBCD)] up to 10 hr with no notable clinical signs. After 3 days washout between dose administration, pups received up to 20 mpk (7.5, 15, and 20 mpk) ALLO by IV infusion, which extended up to 4.8 hr. Dose-dependent sedation was observed at all IV infusion dose levels. After determining that ALLO was well tolerated by extended infusion, bolus IV doses were escalated up to 50 mpk in 1-2 pups (age range PND27-59) per dose level. Hematology and clinical chemistry values were within normal ranges. In summary, ALLO can be administered as IV bolus or extended infusion (>8hr) to conscious neonatal dogs. Pups tolerated ALLO in 20%HPBCD vehicle up to the highest doses tested; 20 mpk (PND17) by infusion and 50 mpk (PND59) by bolus. Conscious neonatal pups appear to tolerate higher doses of ALLO as compared with ALLO administered under isoflurane anesthesia suggesting additive and/or synergistic sedative effects.

1832 Oxidative and Genetic Damage in School Children from a Heavily Polluted City.

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Ecatepec County is one of the most industrialized, populated and polluted counties in Mexico. Particulate matter (PM0.5 and PM2.5) concentrations in this area are frequently above the quality air Mexican standards. Other contaminants, such as lead (Pb) are present. PM and Pb are associated with adverse effects, including oxidative and genetic damage. A cross-sectional study was conducted in 185 schoolchildren aged 7-10 years of both genders from two regions of Ecatepec; an industrialized area with heavy traffic (Zone 1; n=94) and a zone with moderate vehicular traffic (Zone 2; n=91). PM was obtained by air sampling, blood lead (Pb) levels were determined by atomic absorption spectroscopy, plasma malondialdehyde (MDA) as oxidative stress indicator by colorimetry and the DNA damage in mononuclear cells by the comet assay; parents answered a structure questionnaire. PM0.5 and PM2.5 levels, as well as the organic and elemental carbon content (OC and CE, respectively) were higher in Zone 1. Children from this zone showed significantly higher Pb (62% showed >5 μg/dl) and MDA levels, but similar DNA damage compared to Zone 2 children. We observed positive associations between PM0.5 and PM2.5 and OTM values (p<0.05); while a positive association between CO in PM0.5 and PM2.5 and OTM (p<0.05) was also found. MDA levels were associated with PM2.5 and Pb levels, as well as with CO and CE content in both particles (p<0.05). Results suggest that children from Ecatepec are exposed to high PM concentrations and that CO and CE content in PM as well as Pb are involved in the genetic and oxidative damages observed in children living in this highly polluted area. Supported by CONACYT-Mexico (Grant #160534).

1833 Developmental Immunotoxicity Testing for Hazard Identification.

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The inclusion of parameters to assess developmental immunotoxicity (DIT) is an important and critical step forward in regulatory toxicity testing of chemicals. We have performed multiple juvenile and generation studies in rats using various study
designs and test compounds (methylmercury, di-octyl dihloride (DOTC), di(2-ethylhexyl) phthalate, nonylphenol) and the results demonstrate the relatively high sensitivity of immune parameters compared to developmental parameters. Specifically, functional immune parameters appeared affected at relatively low doses. An expanded T cell dependent antibody response (TDAR) parameter set and evaluation of LFS-stimulated NO and TNF-α production by adherent splenocytes were identified as sensitive functional immune parameters. For example, in a juvenile exposure study DOTC affected KLH-induced lymphocyte proliferation at BMD 0.29 mg/kg whereas body weight was affected at 57.7 mg/kg. In a generation study design, alcohol affected splenocyte proliferation at BMD 0.49 % whereas developmental delay was noted at 1.2 %. These results support the OECD TG 445 extended one-generation reproductive toxicity study (OEGRTS) guideline, including its cohort for DIT assessment. It provides substantial insight in the immunotoxic potential of chemical. Our research thus identified complementary immune parameters as potentially useful additions to the EOGRTS guideline which can be easily added to the study protocol. Furthermore, our research demonstrated the relative sensitivity of the juvenile immune system (postnatal day 10-50) and the significance of the juvenile window in DIT testing. The comparisons of various scenarios have provided important lessons about parameter assessment and exposure protocols, which will feed into the definition of a preferred approach to regulatory developmental immunotoxicity testing for chemicals.

Objective

To evaluate the relationship between lung function and arsenic levels in children chronically exposed to arsenic.

Results

At present, 390 children have been included. The mean age is 9.0 years and 98% have lived in the community all their life. 77.6 % of the children were conceived in their current communities. From the studied population, 6.3% reported chronic cough for more than 2 years and 2.9% for 7-12 years. In addition, 12.1% have been treated for bronchiolitis. The mean urinary arsenic level was 155.4 ppb (range of 14.6-893.8). In all subjects spirometric values of FVC, FEV1 and FEV1/FVC ratio were lower with respect to reference values.

Conclusion

A high incidence of lung diseases and a reduction in the lung function were recorded in children chronically exposed to arsenic in drinking water.

1836 Blood and Urine Cadmium Concentrations and Micronucleus Frequency in Buccal Epithelium from Children in Three Populations in Yucatan, Mexico.

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Buccal epithelium represents the first boundary against inhalation or ingestion of toxicants. Micronucleus (MN) tests of buccal epithelium have been effectively used in epidemiological studies in adult populations exposed to genotoxicants; yet, its use in juvenile populations has been limited. A transversal study was conducted in children from the cities: Merida, Progreso and Ticul, in the Yucatan Peninsula to evaluate the MN frequency in exfoliated buccal epithelium and cadmium concentrations in blood and urine from children exposed to different scenarios. Ten children (6 and 8 years of both genders) were selected from each city (n=30). MN frequency was analyzed by microscopy after a Shift staining and Cd concentrations by atomic absorption spectroscopy. MN frequency was similar in the three populations (p=0.7451), Merida 0.33 (0.33/1000 cells), Progreso 0.33 (0.1/1000 cells), and Ticul 0.66 (0.233/1000 cells). Cadmium levels in urine were higher (p=0.043) in children from Merida (State Capital) compared to the other cities (1.30 vs 1.09 and 0.73 ug/l). However, urine Cd concentrations did not correlate with MN frequencies (p=0.505). Blood Cd levels were similar in the three populations (p=0.6021), and a significant and positive correlation between MN frequency and blood Cd levels was observed (rs=0.4583, p=0.0484). Our preliminary results emphasize the importance of conducting biomonitoring of metals and early detection of genotoxic effects in children. Supported by CONACyT-México, Grant # FOSEC-Salud 139738.

1837 Orellanine, a Bipyridyl Mycotoxin, Induces Apoptosis in a Cell Culture Model of Parkinson's Disease.

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Parkinson's disease (PD) affects over one million people in the U.S. alone and more than 6 million people worldwide. Potential risk factors for PD include aging, genetic alterations, and environmental genotoxicant exposures. Although incompletely understood, the etiology and pathological mechanism of PD is characterized by a profound degeneration of dopaminergic neurons in the substantia nigra pars compacta. The majority of PD patients do not know the specific trigger of their disease although exposures to environmental factors are believed to contribute or influence the development and the study of the disease. We have identified a novel environmental bipyridyl mycotoxin similar to Paracut (PQ) and MPTP. Like PQ and MPTP, orellanine (OR) is a bipyridyl molecule and we hypothesized that OR will induce apoptosis in N27 cells. Here, we investigated the neurotoxic effects of OR (3.5,4,4'-tetrathydroxy-2,2'-bipyridine-1,1'-dioxide) on mesencephalic dopaminergic neuronal cell line (N27 cells), an in vitro model of PD. Using an MTT assay, OR induced a dose-dependent decrease in the viability of N27 cells with an EC50 of 44.9 ± 15.3 μM (six times lower than that of PQ or MPP+), suggesting that OR is more potent than PQ or MPP+ on N27 cells. To explore the
mechanisms of cell death, we investigated the effect of OR on mitochondrial-depen-
dent apoptotic pathway in N27 cells. The present study showed that OR induces mitochon-
drial cytochrome C release followed by sequential activation of cas-
pase-9 and caspase-3, and subsequently DNA fragmentation in a dose- and
time-dependent fashion, with peak activation occurring 12 h after OR exposure.
Co-treatment with caspase-3 specific inhibitor, Z-DEVD-FMK (50 mM) signifi-
cantly attenuated OR-induced caspase-3 and DNA fragmentation. Together, this
study demonstrates that OR induces cytotoxicity in mesencephalic N27 dopamine-
producing cells via the caspase-3-dependent apoptotic pathway.

1838 Trichostatin A (TSA), a Histone Deacetylase Inhibitor, Mediates Dopaminergic Cell Death via an NF-κB Dependent Mechanism.
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Trichostatin A (TSA) is a potent, reversible inhibitor of histone deacetylase
(HDACi) that functions through hyperacetylation of core histones. A previous
study from our lab has linked histone hyperacetylation to dopaminergic neurode-
generation. TSA has been shown to induce profound dopaminergic neuronal cell
death, but the cellular mechanisms underlying HDACi-mediated apoptotic cell
death remains unclear. Herein, we show that TSA treatment induced dose depend-
ent apoptotic cell death in the dopaminergic neuronal cell culture model (N27
cells). In order to better define the apoptotic cell death, we investigated the role of
caspases including caspase-2, caspase-8, caspase-9, and caspase-3. The caspases (2, 8, 9, and 3) activation coincided with ROS generation and proteasomal dysfunction
and was found to occur early and prior to cell death. Additionally, TSA induced
apoptotic cell death was preceded by PKC delta activation, upregulation of p44/42
MAP kinase, nuclear translocation of p-65 and hyperacetylation of histone (H3).
Conversely, downregulation of IkBα and survivin levels was also observed as re-
vealed by Western blot and immunohistochemical analyses. Notably, pharmacolog-
ical inhibition of NF-kB and caspases via SN-50 and ZVAD, respectively, conferred
resistance against TSA induced apoptotic cell death in dopaminergic neuronal cells.
These results suggest that NF-kB-mediated and caspase dependent cell death sig-
naling events may be critically linked to TSA induced dopaminergic neuronal de-
generation. Furthermore, the mechanism of the neurotoxicity of the TSA has been
identified extensively. More recently it has been shown that SNpc neurons exhibit L-type
calcium channel, Cav1.3, mediated autonomous pace-making, which could poten-
tially result in increased cytosolic calcium resulting in neurotoxicity. While the VTA
neurons also exhibit pace-making, it is driven by sodium conductances thereby en-
suring that cytosolic calcium levels are not elevated. Thus, Cav1.3 channels could
play a role in the selective susceptibility of SNpc neurons in PD. A short splice vari-
cant of Cav1.3, Cav1.3-42A has been shown to promote calcium influx into the cell
in contrast to the long variant Cav1.3. Thus the presence of greater amounts of
Cav1.3-42A relative to Cav1.3 could lead to increased intracellular calcium. We
measured the relative levels of Cav1.3 and the short splice variant Cav1.3-42A in
brain regions and found that the expression of Cav1.3-42A was 4-fold higher in the
ventral midbrain compared to striatum or cortex. Concomitantly, levels of Cav1.3
were significantly less in ventral midbrain. These results indicate that the presence
of Cav1.3-42A in significantly higher concentration in ventral midbrain, could
contribute not only to amplified pace-making but also result in increased cytosolic
calcium levels in neurons thus contributing to the degeneration of these neurons.

1839 Chemically-Induced Aging of PC12 Cells to Study In Vitro Neurodegeneration.
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Netherlands.
Neurodegenerative diseases, e.g. Parkinson’s disease, are multifactorial and the
mechanisms underlying these disorders are often unknown. In vitro models can in-
crease insight into the cellular and molecular mechanisms by reducing multifactor-
ial diseases to a more controllable set of parameters. A well-known cell model in in
vitro neurotoxicology is the pheochromocytoma (PC12) cell line. PC12 cells have
been used for decades to study the effects of environmental stressors on important
processes of neuronal development, function and degeneration, including different-
iation and neurotogenisis, the synthesis, storage and release of neurotransmitters
and the regulation and function of different ion channels. However, the use of
PC12 cells to investigate neurodegeneration in vitro is debated as this tumor-de-
rivived cell line is rather resistant against environmental insults.
In this study we therefore induced different degrees of aging in PC12 cells by alter-
ing their oxidative status to investigate if this would increase the susceptibility to
environmental stressors, such as pesticides. The characteristics of these aged PC12
subtypes and their sensitivity to an environmental stressor were investigated using
different (functional) assays, i.e. measurements of cell viability, measurements of
oxidative stress using the ROS-sensitive dye H2DCFDA, and single cell measure-
ments of [Ca2+], using the Ca2+-sensitive dye Fura-2.
Our data demonstrate that the different aged PC12 models show a clearly different
Ca2+-homeostasis as compared to naive PC12 cells. This altered homeostasis is ac-
companied by a change in ROS production over time and by an increased sensitiv-
ity to an environmental stressor as compared to naive PC12 cells. As such, these
aged PC12 models with increased vulnerability can be used to gain mechanistic in-
sight in in vitro neurodegeneration studies.
Funding: ZonMW NL (85300003).

1840 Differential Alternate Splicing of L-Type Calcium Channel in
Brain Regions: Implication in Parkinson’s Disease.
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Bangalore, India. Sponsor: B. Moorthy.
Parkinson’s disease (PD) is a movement disorder characterized by resting tremors,
bradykinesia and rigidity caused by death of dopaminergic neurons in substantia
nigra pars compacta (SNpc) of the brain. An intriguing question in the pathogene-
sis of PD is the selective degeneration of SNpc neurons and their terminals in stria-
tum, while the adjacent dopaminergic neurons in ventral tegmental area (VTA) are
affected only in later stages of the disease. Among the several hypotheses that have
been put forth to address cell death, mitochondrial dysfunction, oxidative stress and
accumulation of misfolded proteins in the cytosol (Lewy body) have been stud-
ied extensively. More recently it has been shown that SNpc neurons exhibit L-type
calcium channel, Cav1.3, mediated autonomous pace-making, which could poten-
tially result in increased cytosolic calcium resulting in neurotoxicity. While the VTA
neurons also exhibit pace-making, it is driven by sodium conductances thereby en-
suring that cytosolic calcium levels are not elevated. Thus, Cav1.3 channels could
play a role in the selective susceptibility of SNpc neurons in PD. A short splice vari-
cant of Cav1.3, Cav1.3-42A has been shown to promote calcium influx into the cell
in contrast to the long variant Cav1.3. Thus the presence of greater amounts of
Cav1.3-42A relative to Cav1.3 could lead to increased intracellular calcium. We
measured the relative levels of Cav1.3 and the short splice variant Cav1.3-42A in
brain regions and found that the expression of Cav1.3-42A was 4-fold higher in the
ventral midbrain compared to striatum or cortex. Concomitantly, levels of Cav1.3
were significantly less in ventral midbrain. These results indicate that the presence
of Cav1.3-42A in significantly higher concentration in ventral midbrain, could
contribute not only to amplified pace-making but also result in increased cytosolic
calcium levels in neurons thus contributing to the degeneration of these neurons.

1841 Development of a Cellular Model to Assess Catecholamine
Transport and Toxicity.
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University, Atlanta, GA.
Environmental exposure to neurotoxins can lead to catecholaminergic degenera-
tion and subsequent Parkinson’s disease (PD) development. Historically, dopamine
neurons have been the primary focus of neurotoxicological research in PD though
the importance of noradrenergic degeneration is becoming increasingly evident.
Our laboratory recently developed a cellular model to assess dopaminergic trans-
port and toxicity. Here we demonstrate the development of a novel noradrenergic
cellular model for assessing catecholamine transport and toxicity. Many neurotoxic
compounds act as inhibitors of the vesicular monoamine transporter 2 (VMAT2)
within monoaminergic neurons. The primary role of VMAT2 is to sequester
monoamines into vesicles, protecting them from cytosolic metabolism, and read-
yying them for release upon stimulation. Inhibition of vesicular packaging results in
increased metabolism of neurotransmitter. NE metabolites inhibit mitochondrial
complex I, thereby reducing ATP synthesis and likely inducing toxicity. To investi-
gate noradrenergic transport and toxicity we are utilizing a monoamine-like fluo-
rescent substrate (Molecular Devices) to investigate VMAT2 function in HEK cells
stably transfected with the human norepinephrine transporter (NET) and
mCherry-tagged human VMAT2. These cells model physiological NE uptake
within NE neurons. NET localizes to the cell membrane and VMAT2 forms intra-
cellular vesicle-like structures. The fluorescent substrate we are utilizing mimics the
action of NE in these cells. It is transported into the cell via desipramine-sensitive
NET uptake (IC50 = 113.9 nM). Within the cell, the dye is sequestered into ves-
icles by VMAT2, evidenced by co-localization of fluorescence between the substrate
and mCherry. Tetrabenzene, a VMAT2 inhibitor, inhibits co-localization in a
dose-dependent manner (IC50 = 382.8 nM). Both co-localization and inhibition are
detectable by microscopy and analyzed with IDEV (Cellomics). This cellular model
allows investigation of both normal and altered transporter function in re-
sponse to toxins in a noradrenergic cellular model.

1842 Alpha-Synuclein Protein Aggregates Activate Microglia
and Contribute to Neurotoxicity in the Nigral Dopaminergic
System through a Fyn Kinase Dependent Mechanism.
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Abnormal protein aggregation and chronic neuroinflammation are recognized as
key pathophysiological hallmarks of many neurodegenerative diseases including
Parkinson’s disease (PD). Aggregates of the presynaptic protein α-synuclein are the
1843 Loss of NF-κB p50 Enhances LPS-Induced Systemic Inflammation and Early Microglial Activation.

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Infection associated with chronically activated microglia has been implicated in the progressive degeneration of nigral dopaminergic (DA) neurons in Parkinson’s disease (PD), but the mechanisms are not well understood. NF-kB p50 is an important regulator of the pro-inflammatory response with both inhibitory and initiating roles in the production of cytokines. Importantly, reduced NF-kB p50 expression has been observed in the substantia nigra (SN) of patients suffering dementia with lewy bodies, supporting a potential role for NF-kB p50 in PD-like neurodegeneration. To examine the consequences of loss of NF-kB p50 function in microglial activation in vivo, NF-kB p50+/+ and NF-kB p50−/− mice were injected with 5mg/kg LPS IP and sacrificed after 3 hours. Expression of the pro-inflammatory TNFα, IL-10 and IL-12 cytokines was assessed in unstimulated NF-kB p50+/+ and NF-kB p50−/− microglia. Western blot analyses revealed that Fyn was expressed in primary murine microglia as well as in BV2 microglial cells. Stimulation of the cells with α-synuclein aggregates rapidly activated Fyn within 30 minutes of exposure. The time course activation of Fyn also paralleled PKC α phosphorylation. Interestingly, the α-synuclein induced phosphorylation of PKCα was suppressed in primary microglia isolated from Fyn knockout (Fyn−/−) mice, implicating Fyn as the kinase that carries out this event. The α-synuclein induced activation of α44/42 (Erk1/2) and p38 map kinases was diminished in both Fyn+, as well as PKC8 knockout microglia. Notably, the Fyn knockout primary microglia showed attenuated release of cytokines TNFα, IL-10 and IL-12 in response to α-synuclein aggregate stimulation when compared to wild type microglia. Collectively, our results demonstrate a non receptor tyrosine kinase mediated pro-inflammatory signaling pathway that may mediate neuroinflammation in PD (supported by NIH grants ES19276 and NS65167).

1845 Mutations in pINK-1 and pDR-1 Result in Reduced Dopaminergic Neurodegeneration after Chemical Insult in Caenorhabditis elegans.

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Mitochondrial dysfunction has been linked to neurodegenerative diseases including Parkinson’s disease. Neurons are cells with a high energy demand, and as a result are hypothesized to be particularly vulnerable to disturbances to mitochondrial homeostasis. We investigated how knocking out two genes involved in mitochondrial dynamics, pINK-1 and pDR-1, affected dopaminergic neuron viability after chemical insult in Caenorhabditis elegans. pINK-1 and pDR-1 are the nematode homologs of the human genes PARK1 and PARK2 which when mutated cause familial Parkinson’s disease. We induced neurodegeneration with 6-hydroxydopamine (6-OHDA) and assessed damage 48 hours post-exposure using the BY250 (dat-1::GFP) strain in which the four cephalic dopaminergic neurons are visualized with GFP. We used the BY250 strain as wild-type for pINK-1 and pDR-1. The strain BY250 was generously provided by Michael Aschner (Vanderbilt University) and the strains pINK-1::dat-1; GFP and pDR-1; dat-1::GFP were kindly provided by Guy Caldwell (University of Alabama). Each neuron was scored from 0 to 2.5, with zero representing an intact dendrite and 2.5 representing the highest level of neurodegeneration observed, utilizing fluorescent microscopy. Our preliminary results indicate that after exposure to 15 mM 6-OHDA, the wild-type worm had a higher level of neurodegeneration than the pINK-1 knockout (p<0.0001) and the pDR-1 knockout (p=0.0052). These counterintuitive results might be explained by the fact that loss of pINK-1 promotes mitophagy, which in turn could protect from 6-OHDA damage through mitochondrial functional complementation. Because 6-OHDA generates reactive oxygen species within the cell, it may cause lipid peroxidation and mitochondrial DNA damage, possibilities that we are currently testing. Future studies will explore the effect of these mutations on mitochondrial homeostasis and neurodegeneration after exposure to 6-OHDA and other toxicants associated with Parkinson’s disease including rotenone and paraquat.

1846 Identification of Novel Genes and Epigenetic Mechanisms in C. elegans Models of Idiopathic Parkinson’s Disease and Manganism.

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Background: Idiopathic Parkinson’s disease (PD) and manganism are oxidative stress-related disorders that result in abnormal dopamine (DA) signaling and cell death. Both neurological disorders involve basal ganglia and mitochondrial dysfunction, and suggest overlapping epidemiology, yet the origin of the pathogenesis and the molecular determinants common to both disorders are ill-defined. Recently we have shown that the PD-associated transcription factor SKN-1/Nrf2 is expressed in C. elegans DA neurons and inhibits PD-associated DA neurodegeneration. Aims/Objectives: In this study we asked what are the common genes, molecular pathways, and mechanism involved in DA neuron vulnerability to PD-associated toxicogenic VOGs, we focused on a 30-fold increase in oct-3-ol. Employing the available genetic and molecular tools in our Drosophila model, we demonstrate the modulatory effect of 1-octen-3-ol on dopamine regulatory pathway. We show that 1-octen-3-ol inhibits the vesicular monoamine transporter, VMAT thereby causing the induction of oxidation of dopamine and loss of dopaminergic neurons. Furthermore, we show that 1-octen-3-ol alters regulation of JNK and Akt signaling pathways in flies. In conclusion, our data present strong cues in support of Parkinson-mimetic activity of 1-octen-3-ol and provide insights into neurological problems especially movement disorders associated with damp indoor environments.

1844 Fungal Volatile Organic Compound(s): Potential Environmental Agent(s) for the Pathogenesis of Parkinson’s Disease?

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Parkinson’s disease (PD) is the most common movement disorder and possesses multifactorial etiology. Recently, environmental contaminants including mold exposure are implicated in PD-like movement disorders in epidemiological studies. Although the exact fungal component responsible to such movement disorder is under speculation, fungal volatile organic compounds (VOCs), a class of fungal secondary metabolites has been reported to be associated with the adverse health effects in the human population. There are few in vitro and human studies demonstrating the biological activity of these fungal VOCs; however, the plausible studies elucidating the mechanism of action of these VOCs are lacking. Our lab has pioneered invertebrate Drosophila model to study the neurotoxic effects of common fungal VOCs (Inamdar et al., 2010). We have reported that exposure to chemical standards of 1-octen-3-ol and other fungal VOCs cause loss of dopaminergic neurons and movement deficits in Drosophila suggesting that these tested fungal VOCs have potential to induce Parkinson’s disease like symptoms. To determine the mechanism of toxicity, we focused on 1-octen-3-ol. Employing the available genetic and molecular tools in our Drosophila model, we demonstrate the modulatory effect of 1-octen-3-ol on dopamine regulatory pathway. We show that 1-octen-3-ol inhibits the vesicular monoamine transporter, VMAT thereby causing the induction of oxidation of dopamine and loss of dopaminergic neurons. Furthermore, we show that 1-octen-3-ol alters regulation of JNK and Akt signaling pathways in flies. In conclusion, our data present strong cues in support of Parkinson-mimetic activity of 1-octen-3-ol and provide insights into neurological problems especially movement disorders associated with damp indoor environments.
post-translational modifications of involving these and other proteins identified using RNAi affects DA neuron vulnerability. Conclusions: This study identified novel genes and molecular pathways involved in DA neuron vulnerability in PD and manganism, and provides the first in vivo evidence that a common epigenetic mechanism likely plays a significant role in both disorders. Support: NIEHS ES014459 and ES003299 to RN, and EPA STAR Graduate Fellowship to NVD.

1847 Decreased Mitochondrial Biogenesis and Suppression of Mitochondrial Gene Expression Induced by Environmental Toxins in Caenorhabditis elegans Model of Parkinsonism.

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Mitochondrial alterations have been documented in the brains of subjects with Parkinson’s disease (PD), a disorder characterized by the selective loss of dopamine (DA) neurons. Recent studies demonstrating that PD-associated proteins are either in the mitochondria or translocated into mitochondria in response to stress. Pesticides and heavy metals have been suggested to be risk factors for PD. While, environmental agents can modulate mitochondrial function, the mechanism of this alteration has not been defined in the context of the development and progression of PD. The complexity of the mammalian neurological system has made it difficult to dissect the molecular components involved in the pathogenesis of PD. In the present study we used C. elegans as the model of neuron degeneration and investigated the effect of Mn2+ and rotenone on mitochondrial biogenesis and gene regulation. Exposure to rotenone (2 or 4 μM) resulted in significant loss of dopamine (DA) neuron in C. elegans. We then determined if the rotenone-induced neuron degeneration is accompanied by a change in mitochondria biogenesis. Analysis of mitochondrial genomic replication showed a dramatic decrease in mitochondrial DNA (mtDNA) copies in rotenone-treated C. elegans compared to control. This decreased mitochondrial biogenesis occurred prior to the development of loss of DA neurons, and was persistent. The inhibition of mtDNA replication was also found in C. elegans exposed to neuron toxicant Mn2+ at the concentration 50 or 100 mM. We further examined the mitochondrial gene expression and found significant lower level of mitochondrial complex IV subunits COI and COII in C. elegans exposed to rotenone. These results demonstrated that environmental chemicals cause persistent suppression of mitochondrial biogenesis and mitochondrial gene expression, and suggest a critical role of modifying mitochondrial biogenesis in toxicants-induced neuron degeneration in C. elegans model.

1848 A Genome-Wide RNAi-Based Screen for Enhancers and Suppressors of Manganese Toxicity in Caenorhabditis elegans.

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Environmental or occupational exposure to manganese (Mn) causes a neuropathy resembling idiopathic Parkinson’s disease (PD). Exposure to excessive Mn levels has been linked to mitochondrial dysfunction, oxidative stress, protein aggregation and disruption of iron homeostasis. However, the mechanisms behind these impairments remain unknown, partially due to limited knowledge about genetic factors associated with Mn toxicity. The complexity of the vertebrate brain and the difficulties associated with vertebrate models of PD has hindered the understanding of molecular mechanisms associated with this disorder. The nematode, Caenorhabditis elegans (C. elegans) and mammals share a highly conserved genetic code, and offers an innovative and powerful platform to evaluate the genetic factors associated with Mn-induced toxicity. Taking advantage of the RNA interference (RNAi) technique and the available RNAi library, we are able to evaluate the impact of - 87% of C. elegans genes on Mn toxicity, individually. By feeding the worms with a specific strain of RNAi bacteria, we have successfully knockd down (quantified by qRT-PCR) three selected genes (smf-2, smf-3 and dpy-7) in C. elegans. In vertebrate, the prevalent metal transporter 1 (DMT1) has been shown to regulate Mn transport; its three selected genes (smf-2, smf-3 and dpy-7) in C. elegans.

1849 Loss of Pdr-1/Parkin Alters Manganese Homeostasis in C. elegans.

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Environmental overexposure to the essential trace element manganese (Mn) can result in an irreversible, toxic condition known as manganism. This disorder shares similar neuropathology with Parkinson’s disease (PD), exhibiting overt dopaminergic (D4ergic) cell loss associated with the presentation of motor and cognitive deficits. However, the mechanisms behind the pathophysiology of both disorders remain unclear. Many PD genes have been identified to explain a subset of cases, including the parkin/PARK2 gene that encodes for an E3 ubiquitin ligase. Using Caenorhabditis elegans (C. elegans) as a model that contains the necessary D4ergic machinery, we hypothesize that a loss-of-function mutation in pdr-1, the worm homolog of parkin, would increase vulnerability to Mn toxicity compared to wildtype (WT) worms by altering proper Mn homeostasis. Synchronous L1 worms from WT N2 and pdr-1(ky448) mutant strains were acutely exposed to MnCl2 for 30 minutes, followed by lethality scoring of approximately 40-50 worms 24 hours post-treatment. Here, we show that a loss of pdr-1 increases Mn-induced lethality compared to WT worms (p=0.0005), as seen with a leftward shift in the dose-response curve. Moreover, pdr-1 mutants show higher, dose-dependent Mn accumulation compared to WT worms (p=0.001), suggesting overall impaired Mn homeostasis in pdr-1 mutants. Interestingly, pdr-1 mutants show altered mRNA expression levels of key C. elegans Mn importers, including up-regulation of smf-1 and smf-3, and down-regulation of smf-2. However, pdr-1 mutants do not show any difference in mRNA levels of ferroportin (fep-1) in worms at baseline, indicating a lack of regulation at the export level. Finally, upon exposure, pdr-1 mutants show altered total glutathione (GSH) levels compared to WT animals (p<0.05). Such changes indicate a role of pdr-1 in modulating Mn import through altered transporter expression, implicating a novel role of the PD-associated gene in metal homeostasis and increased oxidative stress.

1850 Manganese Accumulations in Gill Mitochondria of Crassostrea virginica.


Manganese is a neurotoxin causing Manganism in people chronically exposed to elevated levels in their environment. Manganese targets dopamine neurons in basal ganglia. Oxidative stress has been implicated as a factor of manganese toxicity and dopamine dysfunction. Mitochondria play a role as cause and target of oxidative stress damage. The mechanisms of damage is attributed to manganese’s capacity to produce toxic levels of free radicals and induce mitochondrial dysfunction. Other report manganese accumulates in mitochondria and represent the primary pool of manganese in cells. Controversy exists to the extent of manganese accumulation in mitochondria. Others report manganese accumulates within nuclei and cytoplasm, but not mitochondria. Our lab is using the oyster, Crassostrea virginica, as a test animal to study manganese neurotoxicity. We found manganese disrupts the dopamine system as well as mitochondrial respiration. To study if manganese accumulates within mitochondrial of gill cells of C. virginica we used differential centrifugation and atomic absorption spectrometry. Gillis were homogenized and centrifuged to isolate nuclear, mitochondrial and post-mitochondrial fractions. Each fraction was analyzed for manganese. To determine if isolated mitochondrial accumulation we prepared treated mitochondrial suspensions with up to 300 mM manganese. Results show a dose dependent accumulation of manganese in mitochondria of up to 5000%. Two day treatments of animals with 500 and 1000 μM Mn increased manganese (μg/gdw) in gill from a baseline of 5.8 to 41.6 and 133.8, respectively, and centrifugation isolated Manganese, accounting for 87% of total Mn accumulation in nuclear and mitochondrial fractions. This study shows mitochondria accumulate manganese. In vivo treatments reveal accumulations with both the nuclear and mitochondrial fractions.

1851 Manganese Treatments Decreases Immunofluorescence Emissions of Postsynaptic Dopamine D2 Receptors.


Manganese a neurotoxin causing Manganism, a Parkinsons-like disease, disrupts dopamine neurotransmission. Gill lateral cell cilia of Crassostrea virginica are controlled by serotoninergic dopamine D2 receptors. Cilia show cilia-inhibitory effects of dopamine
by blocking dopamine post-synaptic receptors. Questions exist in the literature if manganese decreases the number of D2 receptors in brain. To test that we used antibody-antigen histoimmunofluorescence techniques to visualize dopamine D2 receptors in gill and ganglia of C. virginica. We used a primary antibody against D2 receptors followed by FITC-linked secondary antibody. Animals were treated with 500 μM of manganese for 5 days. Gill, cerebral and visceral ganglia were excised and exposed to primary and secondary antibodies. Paraffin embedded sections were viewed with a phase contrast Zeiss epilume fluorescence microscope with a ProgRes C3 Peltier cooled camera. Antibody treated sections showed bright FITC fluorescence in lateral ciliated cells and other areas of gill and ganglia. Sections lacking primary antibody treatment did not display similar fluorescence. We analyzed fluorescence intensity of 120 control and 80 gill lateral cells from animals treated with manganese using ImageJ software. Intensity of manganese treated cells was 70% less than controls. The study identifies dopamine D2 receptors in gill cells and cerebral and visceral ganglia, and shows a negative correlation between fluorescence intensity of dopamine D2 receptors in manganese treated animals and controls. The question if the decrease in intensity is due to decrease in actual number of receptors or if manganese alters protein conformation of D2 receptor and D2-ligand binding sites needs to be further explored.

Lateral cilia of gill of Crassostrea virginica are controlled by a serotonin-dopaminergic innervation. Dopamine is an inhibitory transmitter at gill causing cilio-inhibition. Manganese is a neurotoxin causing Manganeseism in people exposed to high levels in the atmosphere. Clinical interventions for Manganeseism have not been successful. Recently, p-aminoalicyclic acid (PAS) was reported to provide effective treatment of severe Manganeseism in humans. PAS is an anti-inflammatory drug which has been used to treat tuberculosis. It also has chemotherapeutic properties. Previously, we showed treatments of C. virginica with Mn disrupts dopaminergic innervation of gill. Pre- or co-treatments with PAS or calcium disodium EDTA prevented the neurotoxic effects of manganese. We hypothesized chelating agents would be effective in reversing neurotoxic effects of manganese when applied to gills after manganese. We used gills of C. virginica to measure lateral cilia beating rates of preparations treated first with manganese followed by treatments with either PAS, calcium disodium EDTA or DMSA (meso-2,3-dimercaptosuccinic acid). Beating of cilia were measured by stroboscopic microscopy. Dose responses of PAS, calcium disodium EDTA and DMSA (10⁻⁴ - 10⁻¹ M) against beating were conducted after 100 μM of manganese was added to gill. All 3 drugs reversed the neurotoxic effects of manganese in a dose-dependent manner. DMSA was the most potent. The study demonstrates these chelators are effective in reversing acute neurotoxicity of manganese. This information should be of interest to those designing therapeutic drug treatments for Manganeseism.

Lateral cilia of the gill of Crassostrea virginica are controlled by a serotonin-dopaminergic innervation. Dopamine acts as an excitatory neurotransmitter within the ganglia, but an inhibitory neurotransmitter at gill, causing cilio-inhibition. The mechanism of action of dopamine toxicity is not fully understood, but may be due to oxidative damage. We found several chelators, including p-aminoalicyclic acid (PAS) prevented neurotoxic effects of manganese in C. virginica. The therapeutic actions of PAS are thought to be due to chelation, but PAS is also anti-inflammato-ry. We sought to determine if anti-inflammatory agents and/or antioxidants are effective in preventing neurotoxic actions of manganese in gill of C. virginica. Indomethacin, an anti-inflammatory agent with anti-inflammatory abilities, and ascorbic acid, an antioxidant with possible anti-inflammation abilities were tested. We examined acute and short term (3 - 5 days) treatment of indomethacin and ascorbic acid on manganese toxicity on dopaminergic innervation. Beating rates of lateral cilia in gill epithelial cells were measured by stroboscopic microscopy. Acute or short-term treatments of indomethacin or ascorbic acid (25 - 100 μM) and had no effect on the cilio-inhibitory effects of dopamine (10⁻⁶ - 10⁻³ M). When acute or short-term manganese treated animals (25 - 100 μM) were pretreated with indomethacin or ascorbic acid (25 - 100 μM), both drugs effectively prevented the neurotoxic effects of manganese, with ascorbic acid being more effective than indomethacin. The study demonstrates anti-oxidants and anti-inflammatory agents can block the neurotoxic actions of manganese and may be possible therapeutic agents in the treatment of Manganeseism.

High levels of airborne manganese cause Manganeseism, a neurotoxic, Parkinsons-like disease in humans by interfering with dopaminergic neurotransmission in brain. Recent studies are showing GABAergic neurons also are damaged by man- ganese. C. virginica contains a dopaminergic and serotoninergic innervation of its gill. It is a simple system to study manganese toxicity. Previously we showed manganese disrupts dopaminergic innervation. We also showed the cerebral ganglia of C. virginica contains GABA and within the cerebral ganglia GABA inhibits serotoninergic innervation of gill lateral cell cilia. Here we studied if ganglia and peripheral tissues contain GABA receptors, and if manganese has effects on GABA neurons within cerebral ganglia of C. virginica. We used primary antibodies (GABA and secondary antibodies (IgG-FITC) to detect GABA receptors with paraffinaldehye fixed, paraffin embedded tissues using a Zeiss epilume fluoro- scence microscope and ProgRes C3 Peltier cooled camera. We found fluorescence due to GABA receptors in cerebral ganglia, visceral ganglia, palps and digestive tract. We also examined effects of manganese treatments on GABAergic inhibition of serotonin neurons in cerebral ganglia. Beating of lateral cilia in gill cells were measured by stroboscopic microscopy. Applying serotonin to cerebral ganglia caused a dose-dependent increase in cilia beating. Applying GABA prior to serotonin prevented the increase. Acute applications of manganese (50 and 500 μM) prior to GABA prevented GABA from inhibiting serotonin. The study is showing GABA receptors are present in ganglia and peripheral tissues in C. virginica, and acute manganese treatments damage GABA neurons. C. virginica preparations are good, simple test preparation to study the GABAergic system and the mechanism underlying the neurotoxic effects of manganese.

The advent of high throughput screening (HTS) technology permits identification of compounds that influence various cellular phenotypes. However, screening for small molecule chemical modifiers of neurotoxicants has been limited by the scala-bility of existing phenotyping assays. Furthermore, the adaptation of existing cellular assays to HTS format requires substantial modification of experimental parame-ters and analysis methodology to meet the necessary statistical requirements. Here we describe the successful optimization of the Cellular Fura-2 Manganese Extraction Assay (CFMEA) for HTS. By optimizing cellular density, manganese (Mn) exposure conditions, and extraction parameters, the sensitivity and dynamic range of the fura-2 Mn response was enhanced to permit detection of positive and negative modulators of cellular manganese status. Finally, we quantify and report strategies to control sources of intra- and inter-plate variability by batch level and plate-geometric level analysis. Our goal is to enable HTS with the CFMEA to iden-tify novel modulators of Mn transport. Support: NIH ES016931 and ES010563.

Excessive manganese (Mn) exposure causes a movement disorder commonly referred to as manganeseism in humans. Mn mainly accumulates within mitochondria and adversely affects mitochondrial structure and function both in vivo and in vitro. Over the past decades, mitochondrial dependent biochemical processes such as...
as oxidative stress, Ca2+ dysregulation and apoptotic signaling have been identified as possible mechanisms of Mn neurotoxicity. However, mitochondrial dynamics and the molecular mechanisms that govern the mitochondrial biogenesis during Mn neurotoxicity are yet to be determined. Since the transcriptional co-activator peroxisome proliferator-activated receptor gamma coactivator-1 alpha (PGC-1 alpha) is the master regulator of mitochondrial biogenesis, herein, we examined the effect of Mn on PGC-1 alpha dependent mitochondrial biogenesis in dopaminergic neuronal cells. The MN9D dopaminergic neuronal cell model was exposed to 100-500 μM of manganese, and the mRNA expression levels of several mitochondrial biogenesis markers, including PGC-1 alpha, mitochondrial transcription factor A (TFAM), and cytochrome B (CytB), were measured using qRT-PCR analysis. Interestingly, the results revealed a dose-dependent induction of PGC-1 alpha, TFAM, and CytB mRNA levels following 24 h of manganese exposure, whereas short-time manganese exposure (3-6 h) did not result in any significant induction of PGC-1 alpha mRNA. Since PKD1 serves as a key compensatory anti-apoptotic kinase, we measured PKD1 phosphorylation, and the kinase phosphorylation was significantly decreased after 24 h with 500 μM Mn exposure. A decreased level of Bcl-2 (a pro-survival protein) was also observed at 24 h. Importantly, overexpression of PGC-1 alpha protein significantly protected the cells against Mn-induced neurotoxicity. Taken together, our data indicate that Mn exposure induces mitochondrial biogenesis through PGC-1 alpha transcription to counter metal induced neurotoxic stress (NIH grants ES19267, ES10586, and NS074443).


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Chronic manganese exposure is a well-known occupational and environmental hazard considered to be a potential risk factor for environmentally linked Parkinsonism. α-Synuclein (α-syn) is a major presynaptic protein in CNS, and aggregation of α-syn has been implicated in the pathophysiology of Parkinson’s disease. Previously, we showed that α-syn protects dopaminergic neuronal cells against metal neurotoxicity during early exposure. In the present study, we further characterized the effect of long-term manganese exposure on α-syn metabolism. A 300μM manganese exposure to human α-syn stably expressing N27 dopaminergic cells for 24-48 hr induced a time-dependent increase in α-syn immunoreactive aggregates in the cells. Further analysis of the protein aggregation by inclusion body specific fluorescence probes revealed formation of aggregates in a time-dependent manner. We were also able to detect enhanced accumulation of protein oligomers in manganese treated dopaminergic cells. Studies conducted with MN9D cells stably expressing human α-syn showed that the α-syn is secreted out of the cells into extracellular media following manganese exposure, in a time-dependent manner. Further characterization of extracellular media by ultracentrifugation followed by detection of α-syn by western blotting revealed that manganese treatment induces exosome vesicle formation in α-syn cells. Interestingly, we also observed manganese increased the expression of prion in α-syn expressing cells as compared to vector control cells. Furthermore, manganese treatment increases GRP 78 and caspase 12 levels, suggesting that manganese induces ER stress in α-syn expressing cells. Collectively, these results demonstrate that prolonged manganese exposure promotes α-syn protein aggregation and secretion into extracellular milieu by forming exosomal vesicles, which may contribute to propagation of protein aggregation by a prion-like mechanism in dopaminergic neuronal cells. (NIH grants ES19267, ES10586, and NS075167)

1858 Trp73 Gene in Dopaminergic Neurons Is Highly Susceptible to Manganese Neurotoxicity.

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Chronic exposure to elevated levels of manganese has been linked to a Parkinson’s disease like movement disorder, resulting from dysfunction of the extrapyramidal motor system within the basal ganglia. However, the exact cellular and molecular mechanisms of manganese-induced neurotoxicity remain elusive. Since oxidative stress and apoptosis are considered to be prime mechanisms of manganese neurotoxicity, we sought to identify genes that are susceptible to manganese-induced neurotoxic insult using the Qiang mouse apoptosis RT2 Profiler™ PCR array system. C57 black mice were treated with 10 mg/kg manganese via oral gavage for 30 days. The nigral tissue was collected and RT-PCR based gene expression was performed for 84 genes associated with apoptotic signaling. Interestingly, we found significant down-regulation of a tumor repressor gene, namely Trp73, in manganese treated nigral tissue. In the present study, we further investigated the effect of Mn on Trp73 gene expression following 24 h exposure to elevated levels of Mn in manganese treated N27 dopaminergic cells as well as in manganese treated nigral lysate. Furthermore, manganese treatment in primary striatal culture down-regulated Trp73 protein level in a dose-dependent manner. After confirming Trp73 down-regulation during manganese toxicity in three different models, we have begun additional mechanistic studies in cell culture model of manganese neurotoxicity. Taken together, our results demonstrate Trp73 is a gene susceptible to manganese neurotoxicity, and further characterization of the role of Trp73 in cell survival/cell death will improve our understanding of the molecular underpinnings of manganese neurotoxicity (supported by NIH grants ES10586 and ES19267).

1859 Mechanism of Manganese-Inhibited Induction of Glutamate Transporter GLAST Function.

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Manganese (Mn) is an essential trace element required for normal growth, development and certain coenzymes; however, chronic exposure to elevated levels of Mn induces a Parkinson’s disease-like symptoms referred to as manganism. It has been reported that manganism is associated with excitotoxicity, resulting from the dysfunction of the astroglial glutamate transporters (GLAST and GLT-1), which take up ~80% of synaptic glutamate. However, the mechanism by which Mn disrupts glutamate transporter function has yet to be elucidated. Our previous studies have shown that Mn suppressed transcriptional activity of astrocytic GLAST by decreasing its mRNA and protein levels. Accordingly, herein, we seek to understand the molecular mechanism of Mn-induced GLAST suppression at transcriptional levels, testing whether Mn induces a modulatory effect on astroglial GLAST promoter activity. The experiments were conducted by assessing EAAT1 (human form of GLAST) luciferase activity in astrocytes transiently transfected with EAAT1 vectors. We hypothesized that Ying-Yang 1 (YY1) and NF-kB pathways play a critical role in repressing the GLAST gene expression, and that Mn modulates the activities of these transcriptional factors. The results revealed that Mn activates the YY1 promoter (p<0.05), and overexpression of YY1 decreases EAAT1 promoter activity. We have also found that CBP (CREB binding protein) acts as a co- regulator of YY1 on the EAAT1 promoter. NF-kB also regulates EAAT1 promoter activity. Overexpression of NF-kB p65 increases the EAAT1 promoter activity (10 folds, p<0.001), whereas mutation on NF-kB binding sites at -533 or -163 of EAAT1 promoter represses EAAT1 promoter activity (p<0.001) and reduces p65 effects on EAAT1 promoter activity (3 folds, p<0.001). Mn activates NF-kB reporter activity and produces TNF-α which decreases EAAT1 activity via NF-kB activation. Taken together, our results indicate that the YY1 and NF-kB pathways play critical roles in Mn-induced repression of EAAT1 promoter activity.

1860 Analysis of Brain Mn Distribution Influenced by Disease Stage in Mouse Models of Huntington’s Disease.

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Huntington’s disease (HD) is a genetic neurodegenerative disorder primarily affecting the striatum. There is considerable variability in age of onset in HD strongly influenced by environmental factors. We have previously reported a disease-toxicant interaction between HD and manganese (Mn) exposure, wherein the pathogenic HD mutation is associated with a striatal specific defect in net Mn uptake. For example, there is decreased Mn accumulation in the striatum of FVB-YAC128 mice at 13 weeks of age after 7 day subcutaneous Mn exposure with 3 total injections at 50mg Mn chloride tetrahydrate/kg. To determine if increased exposure duration influenced this phenotype we measured striatal Mn after a 9 day exposure with a total of 5 injections. At this age (13 weeks), prior to the onset of neurodegenerative phenotypes, the higher exposure strengthened the difference in striatal Mn accumulation between wildtype and mutant mice. To determine if this Mn phenotype was progressive over the course of disease, we examined striatal Mn levels at 12 months of age. Unexpectedly, no difference in striatal Mn accumulation between wildtype and mutant mice was observed at 12 months.

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and HD mice was observed using the 7 day paradigm. This alteration in Mn accumulation may be related to disease processes, or to changes due to normal aging, or it may be that the Mn transport defect diminishes with disease progression. To examine this in more detail we will measure and image regional brain Mn accumulation using Inductively Coupled Plasma Mass Spectroscopy (ICPMS), laser ablation ICPMS and other novel methods in HD mouse models. In addition, we will examine primary cultured glia and neurons for cell-type differences in the HD-Mn phenotype. Our ultimate goal is to determine whether disease progression influences brain Mn deposition in HD, and if so, what role glia and neurons play in this process. Supported: NIH ES016931, T32 ES070028

1861 Verification of Manganese-Related Choroid Plexus Differentially Expressed Proteins In Vivo.

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The regulation of brain manganese (Mn) depends largely on the blood-brain barrier and blood-cerebrospinal fluid barrier (BCB). The latter is constituted by choroid plexus (CP) epithelial cells, which is specialized for cerebrospinal fluid (CSF) production, has been considered as a primary target in Mn-induced neurotoxicity. In our previous study, a total of 32 Mn-related differentially expressed proteins were identified by 2D-PAGE combined with Nano-LC-MS/MS in an in vivo toxicity rat model, of which 27 were up-regulated, 5 were down-regulated. In our previous study, a total of 32 Mn-related differentially expressed proteins were identified by 2D-PAGE combined with Nano-LC-MS/MS in an in vivo toxicity rat model, of which 27 were up-regulated, 5 were down-regulated. This study aims to further verify the 7 selected proteins (PHB1, VDAC, β-actin, HSP70, STIP1, TTR, Vimentin) at transcriptional and translational level respectively in immortalized choroidal epithelial Z310 cells in vitro under manganese chloride (MnCl2) exposure. The expressed level of 7 proteins and their mRNA were detected by Western Blot and Real Time RT-PCR, following MnCl2 (0, 50, 100, 200μm) exposure for 24h or 12h in Z310 cells. The results demonstrated that PHB1, β-actin and STIP1 were up-regulated and TTR was down-regulated at both transcriptional and translational levels as compared to controls, which are in accordance with in vivo study. Whereas VDAC, HSP70 and Vimentin were down-regulated at both transcriptional and translational levels as compared to controls, which are opposite to the results in in vivo study. Taking together, this study validated that Mn toxic effects on PHB1, STIP1 and TTR in CP are accurate and reliable, which provide the valuable clue for elucidating the molecular mechanism of Mn toxicity on choroid plexus epithelial cells (partly supported by National Natural Science Foundation of China). Corresponding author: Guojun J. Li, guojun88@yahoo.com.

1862 Expression and Aggregation of a-Synuclein in the Blood-CSF Barrier: New Evidence for the Effects of Toxic Intracellular Manganese and Copper Levels.

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The blood-cerebrospinal fluid barrier (BCB) is responsible for maintaining the homeostasis of a variety of molecules in the brain and cerebrospinal fluid (CSF) including alpha-synuclein (a-Syn), a-Syn plays an integral role in the pathophysiology of Parkinson’s disease. Little is known about the role of the BCB in the transport and regulation of a-Syn in the brain and CSF. Previous findings in this lab provided evidence that a-Syn was endogenously expressed in our immunized Z310 choroidal epithelial cell model, and 100 μM MnCl2 exposure for 24 and 48 hours induced a-Syn aggregation in these cells. The current studies test the hypothesis that the increased a-Syn aggregation in Z310 cells results from Mn interacting with a-Syn expression and/or its aggregation. qPCR was used to quantify a-Syn expression in Z310 cells after 100 μM MnCl2 incubation for 24 hr. Our data revealed that Mn treatment did not affect a-Syn mRNA expression in Z310 cells. Using the Thioflavin T fibril aggregation assay, 70 μM recombinant a-Syn was incubated with 2 μM MnCl2 and CuCl2 for 7 days in vitro. a-Syn aggregation increased significantly (530% and 634%, respectively) as compared to controls (n=3, p<0.05). Finally, we found that 24 hr incubation with 50 and 100 μM CuCl2 induced a-Syn aggregation in Z310 cells. These findings support the hypothesis that cellular aggregation of a-Syn in the BCB is facilitated by exposure to heavy metals. More specifically, Mn exposure induces this effect by two pathways: 1) direct interaction with cellular a-Syn within the cell and/or 2) increasing intracellular Cu levels, shown by our data in literature, leading to Cu-accelerated a-Syn aggregation. (Support by NIH/NIEHS ES08146-S2 Minority Supplemental Award)

1863 Reduced Copper (Cu) Efflux across the Blood-CSF Barrier (BCB) following Manganese Exposure: Effect on Cu Transporters ATP7A and ATP7B.

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Increased Cu levels in blood, saliva and brain are found in Mn-exposed animals and humans. However, the underlying mechanism is unknown. ATP7A and ATP7B are Cu-ATPases that function to maintain intracellular Cu homeostasis by exporting excess Cu from the cytosol to extracellular space. This study was designed to test the hypothesis that Mn exposure disrupted the Cu transport across the BCB by interfering with the intracellular trafficking of ATP7A and ATP7B. Rats received ip injections of 6 mg Mn/kg as MnCl2 or saline, 5 d/wk for 4 wk. Increased Cu and Mn levels in serum and CSF were observed following Mn exposure. An in situ ventricle-cisternal perfusion by using [64Cu] and [14C]-sucrose into brain ventricle was conducted to determine Cu clearance by the BCB. Mn exposure significantly increased [64Cu] radioactivity by 2.6 fold in the CSF outflow, as compared to controls, suggesting a reduced Cu removal by the choroid plexus. Confocal images exhibited both Cu-ATPases distributed in perinuclear region in normal plexus tissues and Z310 cells. Incubation of plexus tissues with Mn or Cu caused translocation of ATP7A from the cytosol toward the apical membrane facing the CSF. Exposing Z310 cells to 100 μM MnCl2 for 24 hr led to a significant decrease in ATP7A and ATP7B fluorescent intensities, which was consistent with their significant mRNA and protein expression reductions in tissue and Z310 cells. The two-chamber Transwell transport studies showed a reduced Cu efflux from the CSF to the blood following Mn exposure or when ATP7A or ATP7B expression were knocked down by siRNA. Collectively, these data suggest that Mn exposure reduces Cu efflux by the BCB which appears to be due to the reduction of ATP7A and ATP7B. A decreased clearance of Cu by Mn exposure may result in the buildup of Cu in the brain. Opposite translocation of ATP7A vs. ATP7B is interesting, yet how this may help interpret a decreased Cu efflux via the BCB remains uncertain. (Supported by NIH/R01-ES080146)

1864 Bone Manganese (Mn) Concentrations in Sprague-Dawley Rats following Subchronic Manganese Exposure.

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Occupational exposure to Mn causes a Parkinson-type disorder called Manganism, a neurodegenerative disease currently without reliable biomarkers for body burden estimation and for pre-symptomatic toxicity assessment. Data in literature suggest that Mn deposited in bone accounts for 43% of total body Mn. The objective of this study was to determine if bone Mn levels were parallel to Mn concentrations in brain regions known to be the targets of Mn toxicity. Groups of rats (6-7 each) received daily dose of 50 mg Mn/kg as MnCl2 PO for 0, 2, 4, 6, 8 or 10 weeks before tissue dissection. The metal concentrations of Mn, zinc (Zn), iron (Fe), and copper (Cu) in bone tissues, blood fluids, brain tissues, as well as other organs were analyzed by atomic absorption spectrophotometry. Following Mn exposure, bone tissues including femur, rib, humerus and skull bones showed a dose-time-dependent increase in Mn concentrations, with 1.0-1.6 μg/g of bone mass at week 10, which were about 2.3-3.6 fold higher than those in controls (~0.4 μg/g at day 0 (p<0.01). A statistically significant relationship exists between bone Mn concentrations and Mn levels in brain tissues such as striatum (r=0.755, p<0.001), hippocampus (r=0.782, p<0.001) and choroid plexus (r=0.652, p<0.001), and in brain fluid such as cerebrospinal fluid (r=0.720, p<0.001). Interestingly, in vivo exposure to Mn also led to significantly increased Fe (152-372%) and Zn (194-230%) concentrations in bone tissues except the skull bones, with no statistically significant effect on bone Cu (58-95%). These results suggest that bone is a significant storage site for body Mn; the good correlation between bone Mn and brain Mn alludes to bone Mn being an internal source of Mn long-term exposure. Further experimentation for noninvasive quantitation of Mn in bone is well warranted for Mn neurotoxicological research. (Supported in part by R21 OH010044, and R01 ES080146)

Keywords: Manganese, Manganism, Biomarker, Bone, Correlation, Pharmacokinetics

1865 Effects of Chronic Manganese Exposure on Cognitive Functioning in Nonhuman Primates.

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Exposure to elevated levels of manganese (Mn) can result in neurological dysfunction in humans including effects on cognitive functioning, particularly those that depend on the integrity of frontal cortex. Our group is performing longitudinal...
studies assessing cognitive and motor function consequent to Mn exposure along with measures of in vivo brain chemistry in non-human primates to better define the neurological consequences and molecular mechanisms of chronic Mn exposure. Adult cynomolgus monkeys were trained to perform cognitive tasks including tests of attention (5 choice serial reaction time (5-CRT)), executive functioning (spatial and non-spatial working memory (self-ordered spatial search (SOSS)) and delayed matching-to-sample (DMTS)), and visuospatial learning (paired associate learning (PAL)). After these (and other) tasks were learned to criterion (using the monkey CANTAB test system) and performance was stable, animals received Mn (1.66 - 2.5 mg Mn/kg per injection, 2x/wk) or vehicle (2x/wk) with an intended total exposure period of 1.5 yrs. Through 8 mos. of exposure, the PAL task, a sensitive index of mild cognitive impairment (MCI) in humans, showed consistent deficits beginning in the second mo. of Mn exposure. Deficits in SOSS and DMTS performance became obvious at 4 and 6 mos. of Mn exposure and attentional impairments, suggested by increased reaction times in the 5-CRT task, were observed consistently at approx. mo. 5. Over the same period of time, performance of the control group either did not change significantly or in some instances, improved. These data show that using a sensitive cognitive assessment battery based on monkey versions of human neuropsychological tests, deficits in several cognitive domains are associated with Mn exposure and in particular, Mn disrupts performance of the PAL task, highly sensitive to detecting early Alzheimer’s disease and MCI, and in particular, disrupts frontal-related functions. Supported by NIH RO1 ES010975.

1866 Neurochemical Alterations in the Nonhuman Primate Brain during Chronic Exposure to Manganese: A 1H MRS Study.

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Alterations in brain chemistry upon chronic manganese (Mn) exposure can be studied non-invasively using 1H Magnetic Resonance Spectroscopy (MRS). In this longitudinal Mn exposure study, 1H MRS was used for the detection of brain metabolite changes that are associated with chronic Mn exposure in non-human primates. A total of 7 adult male cynomolgus monkeys underwent MRS on a 3 T Philips Achieva MRI scanner before Mn exposure and after 8 months of Mn exposure (n=3, 1.66-2.5 mg Mn/kg per injection, 2x/wk). Short echo-time PRESS single voxel MRS data was obtained from frontal cortex (FC), parietal cortex (PC), thalamus, and putamen, and gamma-aminobutyric acid (GABA) edited MRS data was obtained from FC and striatum using MEGA-PRESS. The quantification of metabolites including N-Acetyl-aspartate (NAA), Glutamate (Glu), Glutamine (Gln), nNOS (nNOS-Inositol), total creatine (Cr), choline (Cho) and GABA was done using LCModel. A significant decrease in NAA/Cr (p = 0.035) and mI/Cr (p = 0.013) and a significant increase in Glu/Gln (p = 0.03) from baseline was measured in the PC. The thalamus showed a significant increase in tCr (p < 0.014) and a significant decrease in mH2O (p < 0.022). No significant changes were measured in any metabolites in the FC and in GABA levels in any region. The significant differences in major metabolites are quite robust despite the small number of animals and changes in NAA/Cr in the PC are in agreement with our previously published results (Guilarte et al., Toxicol Sci 94: 351-358, 2006).

1867 What Is the Mechanistic Evidence for Trichloroethylene As a Cause of Parkinson’s Disease?

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Parkinson’s Disease (PD) is a type of movement disorder characterized pathologically, in part, by the progressive and selective loss of dopaminergic neuron cell bodies within the substantia nigra pars compacta (SNpc) and associated deficiency of the neurotransmitter dopamine in the striatum. Mitochondrial dysfunction (i.e., Complex I inhibition), oxidative stress, and abnormal protein aggregation (with intracellular α-synuclein accumulation) have been strongly implicated in PD pathology. A number of recent papers have suggested a causal role for trichloroethylene (TCE) in PD, either directly or via its role in the endogenous formation of 1,1-trichloroethylene-1,2,3,4-tetrahydro-β-carboline (i.e., the TaClo hypothesis). We assess, from a toxicological perspective, the evidence for TCE as a cause or substantial contributor to PD in humans. With regard to the TaClo hypothesis, in vitro studies in a number of different model systems provide evidence that TCE causes changes in mitochondria and dopaminergic neurons but is not preferentially selective towards dopaminergic cells. In vivo studies in rats have shown that high doses of TaClo injected directly into the brain cause modest decreases in dopamine neurons; intraperitoneal (i.p.) administration of TaClo was not selective towards dopaminergic neurons. We observed increases in the striatal concentration of dopamine and its metabolites that were not consistent in direction or with dose. Furthermore, there is no experimental or credible human evidence that TaClo is formed in vivo after exposure to TCE. In animal studies conducted to evaluate TCE directly as a potential cause of PD, relatively high i.p. or oral doses have resulted in findings that are both consistent and inconsistent with mechanistic and motor effects thought to be important in PD. Compared to other well-developed models of PD, the TCE model is poorly characterized. Overall, we conclude that TCE as a causal factor in PD has not been demonstrated and remains speculative.

1868 Neuroprotective Efficacy and Pharmacokinetics of Novel Para-Phenyl Substituted Diindolylmethanes in a Model of Parkinson’s Disease.

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There are no approved therapies that block the chronic inflammatory component of neurodegenerative diseases such as Parkinson’s. This is partly because of poor distribution to the central nervous system for compounds with demonstrated efficacy in vitro. This study examined selected para-phenyl substituted diindolylmethanes (C-DIM) compounds, which we previously demonstrated to be effective at decreasing glial-derived inflammation, gene expression in vitro. We postulated that the pharmacokinetic properties of C-DIM compounds would positively correlate with neuroprotective efficacy in a progressive model of Parkinson’s disease (PD) in vivo. Pharmacokinetics and metabolism of 1,1-bis(5-indolyl)-1-(p-methoxyphenyl)methane (C-DIM5), 1,1-bis(5-indolyl)-1-(p-hydroxyphenyl)methane (C-DIM8), and 1,1-bis(5-indolyl)-1-(p-chlorophenyl)methane (C-DIM12) were determined in plasma and brain of C57Bl/6 mice. Intravenous (1 mg/kg) and oral (10 mg/kg) doses were given to determine the optimal route of administration and putative metabolites were measured in plasma, liver, and urine. Oral dosage of C-DIM compounds displayed greater AUC, Cmax, and Tmax levels than intravenous administration. C-DIM12 exhibited distinguished pharmacokinetics of the selected C-DIMs, with an oral bioavailability of 42% in comparison of C-DIM8 (6%). Following pharmacokinetic studies, efficacy of C-DIM5, C-DIM8, and C-DIM12 (50 mg/kg, oral gavage) was established using a progressive, neuroninflammatory PD model employing MPTP and probenecid (MPTPp) over a period of 14 days. By first creating a lesion of the substantia nigra pars compacta (SNpc) in a number of different cell systems have provided evidence that TaClo causes inflammation in the central nervous system can produce large amount of pro-inflammatory factors such as tumor necrosis factor α, interleukin-1β, interleukin-6, proteins, and glial activation; the subsequent interaction among these factors and host cells leads to progressive death of dopaminergic neurons in SNpc, resulting in PD. Lipopolysaccharide (LPS) is an endotoxin derived from gram-negative bacteria and is widely used as an inflammation inducer. In this preliminary study, we wanted to know what dose level of LPS can effectively induce PD in C57Bl/6 mice by direct brain injection at the SNpc site. Six groups of mice were injected with low dose LPS (200 microgram/Kg), high dose LPS (400microgram/Kg) and normal saline as control, respectively, into one side of the SNpc site or the target it projects to (striatum of the same side). The surviving DA neurons in the SNpc site were shown and counted by immunohistochemistry method using rabbit anti-tyrosine

1869 Neurotoxicity Study of Lipopolysaccharide (LPS) Induced Mouse Model of Parkinson’s Disease.

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Parkinson’s disease (PD) is a debilitating neurodegenerative disease characterized by gradual and progressive loss of dopaminergic (DA) neurons in substantia nigra pars compacta (SNpc). 95% of PD may be attributed to multi-variant etiology; among them, inflammation is getting more attention. Current understanding is that inflammation in the central nervous system can produce large amount of pro-inflammatory factors such as tumor necrosis factor α, interleukin-1β, interleukin-6, proteins, and glial activation; the subsequent interaction among these factors and host cells leads to progressive death of dopaminergic neurons in SNpc, resulting in PD. Lipopolysaccharide (LPS) is an endotoxin derived from gram-negative bacteria and is widely used as an inflammation inducer. In this preliminary study, we wanted to know what dose level of LPS can effectively induce PD in C57Bl/6 mice by direct brain injection at the SNpc site. Six groups of mice were injected with low dose LPS (200 microgram/Kg), high dose LPS (400microgram/Kg) and normal saline as control, respectively, into one side of the SNpc site or the target it projects to (striatum of the same side). The surviving DA neurons in the SNpc site were shown and counted by immunohistochemistry method using rabbit anti-tyrosine
hydroxylase and visualized by diaminobenzidine staining at 1st and 3rd week after injection. We compared the number of DA neurons at SNc with DA neurons at SNc on the contralateral side. We found significant loss of DA neurons at SNc in both injection sites of SNc and striatum in the high dose groups at 1st and 3rd week. No significant loss of DA neurons was observed in the low dose and the control groups. We concluded that LPS at the dose level of 400microgram/kg can effectively and rapidly induce PD in C57Bl/6 mice through direct brain injection, and the pathological change resembles the neurodegenerative characteristics found in PD. The dose response relationship and the pro-inflammatory characteristics associated with the DA neuron loss may need further investigation.

1870 Neuroinflammation and Microglial Dysfunction.

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Neuro-inflammation and accumulation of Ap-containing amyloid plaques are critical components in the pathogenesis of Alzheimer’s disease (AD). Microglia are the brain tissue macrophages that play critical roles in the inflammatory aspects of AD by releasing proinflammatory cytokines. Activated microglia are also able to migrate to the site of inflammation and phagocytose the peritoneal macrophages’ functions. Therefore, we hypothesized that HMGB1 contributes to microglial dysfunction in neuro-inflammation. In this study we demonstrate that HMGB1 levels were significantly elevated in the extracellular space of cultured BV2 macroglia cells 24 hours after exposure to 1 μg/ml Lipopolysaccharide (LPS) compare to untreated control cells. Exposure of BV2 cells to LPS resulted in significant inhibition of microglia’s ability to migrate and to phagocytose. Simultaneous treatment of BV2 cells with LPS and anti-HMGB1 antibody significantly improved the migration and phagocytic ability of these cells. Moreover, treating BV2 cells with recombinant HMGB1 not only induced impairment in migration and phagocytosis but also accompanied by the expression of Toll-like receptor 4 (TLR4) on these cells. These results suggest that activation of the LPS-induced HMGB1/TLR4 signaling pathway contributes to the microglia dysfunction. Thus, inhibiting of HMGB1 may provide a therapeutic target for enhancing of microglia’s ability to migrate and phagocytose in AD.

1871 Translocator Protein 18 KDA (TSPO) in Sandhoff Disease: An Update on a Preclinical Biomarker of Neurodegeneration.

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Translocator protein 18 kDa (TSPO) is extensively used as a biomarker of brain disease and neuroinflammation (Chen & Guillart, Pharm Ther 118: 1-17, 2008). Sandhoff disease, which is clinically similar to Tay-Sachs disease, is a neurodegenerative condition in which a deficiency in the enzyme lysosomal β-hexosaminidase leads to accumulation of gangliosides and glycolipids in the brain, resulting in progressive and widespread neurodegeneration. We previously reported the longitudinal expression of TSPO in a mouse model of Sandhoff disease at the late stages of the disease (2 and 3 months). Now we report TSPO expression at early disease time points (1 and 1.5 months) in order to assess the temporal expression of TSPO and its relationship to behavioral and neuropathological endpoints. Using TSPO quantitative autoradiography with the TSPO specific ligands [125I]iodo-DPA-713 and [3H]-DPA-713, we show that TSPO levels are increased at 1 month and remain high in brain regions corresponding with behavioral expression of disease. We also demonstrate that the temporal increase in TSPO levels is associated with ongoing neurodegenerative changes and activation of microglia and astrocytes using silver staining (marker of active degeneration) and immunohistochemistry of Mac-1 (microglia) and GFAP (astrocytes). Triple labeled immunofluorescent and immunohistochemical imaging confirmed that TSPO colocalized with microglial markers as well as with the gp91phox subunit of NADPH oxidase at an age when brain tissue is undergoing neurodegeneration. These results further strengthen the evidence that TSPO can be used as an easy and sensitive preclinical biomarker of brain injury and inflammation [Supported by NIEHS’s grant ES007062 to TRG].

1872 Characterization of Mice with Overexpression of the Vesicular Monoamine Transporter 2 (VMAT2; S1c182a).

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The vesicular monoamine transporter 2 (VMAT2) is responsible for packaging monoamines for rapid release at neuronal synapses. VMAT2 is also responsible for sequestering toxins away from their sites of action in the cell. Despite previous evidence suggesting that decreased levels of VMAT2 can contribute to cellular dysfunction, it was not known if an increase in VMAT2 protein would result in increased function in vivo. We generated C57Bl6 BAC transgenic mice with increased VMAT2 expression to determine the effects of increased vesicular monoamine filling on associated neurochemical and behavioral outcomes. Our VMAT2-HI mice show robust VMAT2 overexpression, proper localization of VMAT2, increased whole brain synaptic vesicle [3H]-dopamine uptake, and a measurable behavioral phenotype. These data show the effects of elevated VMAT2 levels on neurochemistry and behavior, suggesting that increased vesicular dopamine storage mediated by increased VMAT2 levels can alter the neurochemical and behavioral output of the monoaminergic system. Finally, as VMAT2 acts as a neuroprotective mechanism in monoaminergic neurons, these VMAT2-HI mice serve as a valuable tool to investigate resistance to toxic exposure.

1873 Background Survival of Transgenic Mouse Models of Alzheimer’s Disease in Our Laboratories.


Transgenic mouse models of Alzheimer’s disease are of great value for evaluating efficacy and/or toxicity of drug therapies. These models typically present hallmarkstions of the disease between 12 and 20 months of age; however, age-related attrition presents a challenge that may impair the statistical significance of the data generated. We present survival data of three different transgenic Alzheimer mouse models when housed under our optimized laboratory conditions. The homoygous strain of PDAPP mouse (STOCK-TgN(APP)76-2175) showed mortality rates of 13% (males) and 21% (females), between the ages of 7 to 12 months and of 70% (males) and 60% (females), between the ages of 12 to 21 months. Heterozygous PDAPP mice (STOCK-TgN(APP)76-2175) showed greater survival with only 6.4% and 16.8% mortality for males and females, respectively at 14 to 20 months of age and 16% and 42% mortality for males and females, respectively, between 20 to 26 months of age. Young Tg2576 mice (B6; SJL-Tg (APPSWE) 2576Kha) presented a survival rate comparable to wild type mice with 100% and 90% survival between the ages of 9 and 14 weeks. When survival is calculated for older Tg2576 mice, 79 and 75 % of males and females, respectively, survive over the age of 13 to 22 months. As the survival rates vary between the different strains and the age-related attrition is relatively high, consideration for increased population should be given accordingly in study designs using these transgenic mouse models.

1874 Oxidative Damage and Age-Related Alterations in Kainic Acid-Induced Excitotoxicity.

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Recent research findings in brain have highlighted increased excitatory stimulation as a contributor to aging as well as neuronal damage that accompanies multiple neurodegenerative diseases. Findings from patients and animal models have widely supported the hypothesis that neuronal oxidative/nitrosative damage is a major effect on contributing to neurodegeneration. Therefore, the present study investigated antioxidative and neuroprotective effects of the antioxidant vitamin E (α-toco- pherol) and the NMDA receptor antagonist memantine in age-related excitotoxicity induced by kainic acid (KA). Mice exposed to KA (1 nmol/5 μl, i.c.v.) showed significant increases in cerebral oxidative biomarker F2-isoprostanes (F2-Isopx, 158%) and nitrosative biomarker citrulline (249%) formation when determined at 30 min after exposure. At the same time, pyramidal neurons in the hippocampus of young and old mice showed significant reductions in dendritic length (60%) and spine density (40%) compared to controls (100%). Pretreatment with vitamin E (100 mg/kg, ip/day for 3 days) and memantine (5 mg/kg, ip, but not 1 mg/kg) attenuated the KA-induced increases in cerebral F2-Isopx and citrulline and decrease in...
spine density of hippocampal pyramidal neurons in young mice. However, vitamin E (100 mg/kg, ip/day for 3 days) and memantine (5 mg/kg, ip) were not effective in suppressing KA-induced oxidative stress and a decrease in the dendritic length of hippocampal pyramidal neurons in aged mice. These data strongly suggest that different mechanisms are involved in cerebral neuroprotection of aged mice compared to young mice. Elucidation of these mechanistic changes has important clinical implications for therapeutic strategies in both normal aging and neurodegenerative disease.

1875 Rapid Immunoassay Development and Evaluation of ICAM-1 and E-Selectin As Potential Biomarkers of Vascular Injury in Rats.

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To qualify biomarkers of vascular injury in support of the Predictive Safety Testing Consortium’s (PSTC) Vascular Injury Working Group (VIWG) we developed bio-marker immunoassays on the Glymax platform for rat ICAM-1 and E-Selectin using a fit-for-purpose approach. ICAM-1 (Intercellular Adhesion Molecule 1), also known as CD54, is a cell surface glycoprotein typically expressed on endothelial cells. E-Selectin, also known as CD62 and endothelial leukocyte adhesion molecule 1, is typically expressed on activated endothelial cells. The assays were developed using commercially sourced reagents. Rats (n=5/group) were treated with either 60 mg/kg of CI-1044 (oral gavage) or 100 mg/kg of fenoldopam (subcutaneous). Both drugs are known to cause vascular injury. Samples were taken at 4, 8, 16, 24, 48, 72 hours and recovery for CI-1044 and at 0.5, 2, 6, 24, 72 hours and recovery for fenoldopam and serum was evaluated for levels of ICAM-1 and E-Selectin. Histopathological examination indicated vascular lesions were present from the CI-1044 rats starting at 16 hours post dose consisting of perivascular inflammation of the mesenteric arteries and progressing to fibrinoid necrosis at later time points. The mesenteric vascular lesions in fenoldopam-treated rats consisted of arteriolar necrosis/apoptosis observed at 24 hours post dose, followed by perivascular inflammation and fibrinoid necrosis at later time points. There was no significant change noted in the circulating levels of ICAM-1 in CI-1044 or fenoldopam treatment groups. There was a significant decrease in serum E-Selectin levels in both CI-1044 and fenoldopam treated groups after 48 hours. E-Selectin has the potential to become a valuable biomarker in the evaluation of vascular injury for preclinical drug safety studies.

1876 Synthesis and Characterization of PEG-Ylated Meso-Porphyrins for Targeting the Epidermal Growth Factor Receptor in CRC.


This project introduces a novel macrocycle conjugated to polyethylene glycol linker, which we hypothesize will serve as the template for a selective molecule with high fluorescence yields that greatly enhances earlier detection of CRC by utilizing the EGF as a biomarker. Our targeted remedy is a porphyrin that is conjugated to a peptide with an affinity for the Epidermal Growth Factor Receptor (EGFR). Porphyrins are characteristically aromatically stable, contain trademark absorption bands in the visible and near-IR range, and have fluorescence quantum yields much above the current fluorophores. This makes the macrocycle optimal for confocal laser endomicroscopy (CLE) agent production. Consequently, we use a polyethylene glycol linker in order to increase water solubility, retain low toxicity, and to achieve high fluorescence quantum yields, as well as high conjugation yields. In this research, we were able to produce both precursors to the porphyrin-peptide conjugate, MesopOR-(mono)-3PEG and MesopOR-(di)-3PEG. These molecules were synthesized successfully with the use of peptide conjugation mechanisms. Molecular weights were confirmed using Matrix Assisted Laser Desorption Ionization Mass Spectrometry (MALDI-MS). Characterization was performed using 1H Nuclear Magnetic Resonance (1H-NMR) and Ultraviolet-Visible Spectrophotometry (UV-Vis) of the intended molecules. The synthesized molecules, MesopOR-(mono) 3PEG and MesopOR-(di)-3PEG, will be useful in peptide conjugation that targets EGFR. These peptide ligands will increase selectivity and detect CRC via CLE.

1877 Multiplex Analysis of Urinary Protein Biomarkers for the Detection of Vancomycin Induced Sub-Acute Nephrotoxicity.

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In pharmaceutical and chemical industries the kidney is routinely assessed during preclinical safety evaluations. The importance of the kidney as a central detoxification organ leads to a high exposure of renal tissue to drugs, reactive metabolites or environmental compounds. Traditional markers for assessing renal toxicity, such as blood urea nitrogen (BUN) and serum creatinine (SCr), are insensitive. Although both are direct measurements of renal function, increases in serum concentrations of these biomarkers occur only after substantial renal injury. For improved detection of acute nephrotoxicity a panel of novel urinary kidney biomarkers has been approved by the FDA, EMA and PMDA. However, limited data regarding the performance of these acute markers after sub-acute or sub-chronic treatment are publicly available. To increase the applicability of these markers, it is important to evaluate the ability to detect these markers after 28 days or even longer.

In this study, Wistar rats were treated with three doses of Vancomycin to induce renal damage and studied for 28 days. Urine was collected under cooled conditions on an 18-hour cycle on days 6, 15 and 29. Lumines®-MAP® based MILLI-PLEX® Rat Kidney Toxicity Magnetic Bead Assays were used to measure 14 candidate protein biomarkers simultaneously from the urine samples. Vancomycin treatment resulted in a dose-dependent increase in urinary biomarkers, specific for the observed areas within the nephron, determined histopathologically. Several biomarkers were found promising in this study, which includes NGAL, Cystatin C, KIM-1, Osteopontin, Clusterin and Albumin. The simultaneous measurement of these proteins with multiplex technology offered a robust and convenient method to study these biomarkers. Taken together, our data demonstrate the high accuracy and predictivity of some of these new makers to detect sub-acute treatment with one well described nephrotoxin, Vancomycin.

1878 Association of the Cumulative Body Burden of Estrogen-3,4-Quinone with Body Mass Index and Breast Cancer Risk Using Albumin Adducts As Biomarkers.

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Both 17β-estradiol-2,3-quinone (E2-2,3-Q) and 17β-estradiol-3,4-quinone (E2-3,4-Q) are reactive metabolites of estrogen. Elevation of E2-3,4-Q to E2-2,3-Q ratio is thought to be an important indicator of estrogen-induced carcinogenesis. Our current study compared the cumulative body burden of these estrogen quinones in serum samples taken from Taiwanese women with breast cancer (n=152) vs healthy controls (n=140) by using albumin (Alb) adducts as biomarkers. Results clearly demonstrated the presence of cysteinyl adducts of E2-2,3-Q-4-S-Alb and E2-3,4-Q-2-S-Alb in all study population at levels ranging from 61.7-1330 and 66.6-1590 pmol/g, respectively. Correlation coefficient between E2-2,3-Q-4-S-Alb and E2-3,4-Q-2-S-Alb was 0.610 for controls and 0.767 for breast cancer patients (p<0.001). We also noticed that in subjects under age 50 with body mass index (BMI) less than 27, background levels of E2-3,4-Q-2-S-Alb was inversely proportional to BMI with about 25% increase in E2-3,4-Q-2-S-Alb per 5 kg/m2 decrease in BMI (p<0.001). In addition, we confirmed that mean levels of E2-3,4-Q-2-S-Alb in breast cancer patients were ~5-fold greater than in those of controls (p<0.001). Overall, this evidence suggests that disparity in estrogen disposition and the subsequent elevation of cumulative body burden of E2-3,4-Q may play a role in the development of breast cancer. (This work was supported by the National Science Council, Taiwan, through Grants NSC99-2314-B-005-001-MY3)

1879 Biomarker-Based Evaluation of Diesel Exhaust Emissions from 2007-Compliant Engines in Rats and Mice Exposed for Defined Time Periods.

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In 2001 the USEPA adopted new air quality standards for diesel fuels and emissions; the health impact was not established. Diesel exhaust (DE) is associated with adverse health effects, including cardiovascular, lung cancer, and neurological effects.
Results: PBA were measured by LC/MS in rats dosed rifampicin (a BSEP inhibitor), methylnitroso-diethylammonium (a hepatic toxicant), methylnitroso-phenylendiamine (a muscular toxicant) and dexamethasone (a hepatic glycogenesis inducer). Increase of cholic acid and taurocholic acid, accompanied by decrease of glycocholic acid and chenodeoxycholic acid were specific to rifampicin treatment, confirming the reproducibility of the previous analyses. Next, single dosing studies were conducted in rats and dogs to investigate the time-course changes of PBA. In rats, PBA were increased maximally at 6~9 hrs after cyclosporin A dosing and tended to return to the normal level at 24 hrs after dosing, which corresponded to the toxicokinetics of cyclosporin A. On the other hand, PBA were detected only after feeding in untreated dogs, and rifampicin intensified the rise in PBA concentration induced by meal. [Conclusion] The results show that PBA can be specific BMs of BSEP inhibition in preclinical animals. Since PBA showed time-course changes associated with plasma drug concentrations and were also affected by meal in dogs, plasma sampling time point would be critical to evaluate changes in PBA induced by BSEP inhibition.

1882 Kidney Injury Biomarkers Lipocardin-2, Clusterin and Albumin, but Not Kidney Injury Molecule 1, Are More Sensitive Than Traditional Markers of Gentamicin-Induced Kidney Injury in Beagles.


Safety Assessment, AstraZeneca Pharmaceuticals, Witham, MA; 2Myriad RBM, Saranac Lake, NY; 7Pfizer, San Diego, CA; 8Gala&SmithKline, King of Prussia, PA; 9Janssen Pharmaceuticals, La Jolla, CA.

A panel of 8 novel urinary protein biomarkers were recently qualified by the FDA for the detection of acute kidney injury (AKI), significantly improving the sensitivity and specificity with which renal injury may be detected in rats. However, the translation of these biomarkers to dogs, one of the most common species used in preclinical drug development, remains unknown yet must be established prior to broad implementation in toxicity screens. Here, we measured AKI biomarkers kidney injury molecule 1 (Kim-1), lipocardin-2 (NLGAL), clusterin and albumin in urine from male beagles infused once daily for 10 days with the nephrotoxic antibiotic gentamicin. Histological examination of the kidneys following treatment revealed severe epithelial degeneration, necrosis and regeneration alongside tubular dilatation in gentamicin-treated dogs only. In gentamicin-treated dogs, significantly elevated urinary protein levels of albumin, clusterin and NLGAL were measurable as early as 3 days after start of treatment with peak levels >100-fold above background. Both the speed and magnitude of response detected with these new biomarkers were superior to traditional AKI biomarkers, blood urea nitrogen (BUN) and serum creatinine (sCr), which did not increase until day 8. Urinary Kim-1 levels peaked around 2-fold, an order of magnitude lower than published levels from a similar 10-day rat study. Further analysis of samples for Kim-1 mRNA and protein will tell whether the canine Kim-1 response is markedly different from that observed in rats. These data indicate that three of four novel biomarkers qualified in rat (NLGAL, clusterin and albumin) translate to canine and represent a substantial improvement over traditional AKI biomarkers in preclinical drug development.

1883 Bio-Monitoring of Multimycotoxin Exposure in Human Urine Samples from Cameroon Using a Multibiomarker Approach.

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Mycotoxins are secondary metabolites of fungi that contaminate food, but bio-monitoring of human exposure has mostly been limited to a few individually measured mycotoxins. As the aim of this study was to determine frequency of exposure and level of multi-mycotoxin in urine samples obtained from Cameroonian adults, mostly from HIV infected individuals, 175 urine samples were collected for measurements of cTnT and hematological parameters. Preliminary results showed a release of cTnT in plasma in 8% of mice at 6 mg/kg, 17% of mice at 9 and 12 mg/kg, and 92% of mice at 18 and 24 mg/kg cumulative DOX doses, indicating a dose-related increase in cardiac injury. Plasma cTnT levels were also elevated in 58%, and 55% of mice treated with DXZ+DOX for 3, 6, and 8 weeks, respectively. Microscopic examination of hearts revealed the presence of cardiac lesions only in mice exposed to 24 mg/kg cumulative DOX dose, suggesting that DXZ provided cardioprotection against irreversible damage. The presence of histological damage in the heart of only mice exposed to the highest cumulative DOX dose yet some elevation in cTnT at lower cumulative DOX doses suggests that our ongoing evaluation of molecular changes may reveal more sensitive biomarkers of cardiac injury.

1880 Investigation of Early Biomarkers of Doxorubicin (DOX)-Induced Cardiac Tissue Injury in B6C3F1 Mice.

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Cardiac troponins (cTnT) are widely used as biomarkers for assessing cardiac injury in preclinical and clinic testing. Here we report a study that was designed to identify early biomarkers of cardiac injury in male B6C3F1 mice. Mice were treated weekly for 2, 3, 4, 6, and 8 weeks with (a) 3 mg/kg DOX given intravenously (i.v.) via tail vein, resulting in cumulative doses of 6, 9, 12, 18, and 24 mg/kg, respectively, (b) an equivalent volume of saline (SAL) given i.v. via tail vein, (c-d) 60 mg/kg dexrazoxane (DXZ), a cardioprotective drug, given intraperitoneal (i.p.) 30 min before i.v. injection of DOX or SAL, and (e) SAL given i.p. 30 min before i.v. injection of DOX. At necropsy, which was performed a week after the last dose, blood samples were collected for measurements of cTnT and hematological parameters. Also, hearts were collected for evaluations by light and electron microscopy, genomics, proteomics, and metabolomics. Preliminary results showed a release of cTnT in plasma in 8% of mice at 6 mg/kg, 17% of mice at 9 and 12 mg/kg, and 92% of mice at 18 and 24 mg/kg cumulative DOX doses, indicating a dose related increase in cardiac injury. Plasma cTnT levels were also elevated in 58%, and 55% of mice treated with DXZ+DOX for 3, 6, and 8 weeks, respectively. Microscopic examination of hearts revealed the presence of cardiac lesions only in mice exposed to 24 mg/kg cumulative DOX dose, suggesting that DXZ provided cardioprotection against irreversible damage. The presence of histological damage in the heart of only mice exposed to the highest cumulative DOX dose yet some elevation in cTnT at lower cumulative DOX doses suggests that our ongoing evaluation of molecular changes may reveal more sensitive biomarkers of cardiac injury.

1881 Investigation on Plasma Bile Acids As Biomarkers of Cholestasis Induced by Bsep Inhibition in Preclinical Animals.

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[Purpose] The bile salt export pump (BSEP) is a bile acid efflux transporter, the inhibition of which has been proposed to play a role in drug-induced cholestasis. However, there are no known biomarkers (BMs) to detect BSEP inhibition. We previously performed metabolomics analyses on rat plasma treated with cyclosporin A, a well-known BSEP inhibitor, and found that plasma bile acids (PBA) could be BMs for BSEP inhibition. To further evaluate the usefulness of PBA, we characterized the specificity and time-course changes of PBA in rats and dogs. [Methods and
individualy or in combinations, in 110/175 (63%) samples; from HIV positive (64%) and HIV negative (57%), with additional 4 analytes present only in HIV positive samples. AFM1 (10%); mean 0.5, range <LOQ-1.4μg/L) and FB1 (3%; mean 0.6, range 0.5-15μg/L) were detected in the HIV subpopulation whilst low levels (<LOQ) were found in one sample each from HIV negative group. One HIV positive individual's urine contained 6 metabolites. Levels of these metabolites were generally similar to those reported elsewhere in Africa. For the first time in Africa and elsewhere, this study has reported on 11 mycotoxin biomarkers/bio-measures quantified in human urine. Mycotoxin exposures in HIV individuals may require particular attention. The findings may constitute a major step towards mycotoxin exposure assessment and national mycotoxin regulations in Cameroon.

1884 Development of a New Oxidative Stress Biomarker Dityrosine ELISA.
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Accumulating evidence indicates that oxidative stress plays an important role in various diseases such as cancer, diabetes and hypertension. Recently it is also reported that oxidative stress is involved in toxicity of chemical substances such as arsenic, asbestos, diesel exhaust micro particles and antiinflammatory drugs, and monitoring of oxidative stress inside human body may be informative for toxicological study. Oxidative stress may cause oxidative damages to biomolecules such as nucleic acids, lipids, proteins and enzymes, and oxidized products of such biomolecules have been used for the assessment of oxidative stress in the living bodies. Although protein oxidation is one of the most important biomolecules, only limited number of reports about the oxidized proteins has been published. Tyrosine is one of the major targets of protein oxidation, and dityrosine is known to be formed by oxidative stress. In this presentation, development of a new dityrosine ELISA is reported. A competitive dityrosine ELISA is established using anti-dityrosine monoclonal antibody (clone 1C3) which was developed by Kato et.al. 50 ul of diluted samples or standards are poured into micro plate wells which is pre-coated by dityrosine antigen, and incubated at 4 degree C overnight. After washing by PBS-tween buffer, 100 ul of HRP-conjugated anti mouse antibody is poured, and incubated for 1 hour at room temperature. TMB is used for color development. Assay range of dityrosine ELISA is 0.05 to 12 umol/L, and shows good linearity and reproducibility. Dityrosine concentration in human urine ranges between 0.12 and 3.95 umol/L (mean 1.44 umol/L), and urinary dityrosine concentration measured by ELISA significantly correlated with that measured by LC-MS/MS. In conclusion, dityrosine ELISA may be useful for the assessment of oxidative stress in the living bodies.

1885 Method Development of Serum Canine Inhibin B Enzyme-Linked Immunosorbent Assay (ELISA).
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Inhibin B (INH-B) is a heterodimeric glycoprotein consisting of an alpha and a beta-B subunit linked by disulphide bridges. INH-B is produced by the testes as well as the ovaries, and is responsible for the selective negative feedback control of follicle stimulating hormone. In males, INH-B is synthesized by the Sertoli cells in the testis, and can be used as a marker of Sertoli cell function and spermatogenesis in adult males. Hence, it is being considered a biomarker for detecting testicular damage. INH-B has been quantified in humans, rats and non-human primates, but not in canines due to lack of availability of reagents. Here, we report the methods development of the canine INH-B ELISA from Cusabio Biotech Co. (Wuhan, China). Assay standard curve is ranged from 4 to 1000 pg/ml with serum requirement of 50 μL. Assay optimization included modification of the procedure to include sample mixing followed by prolonged primary antibody-antigen incubation time to ensure saturation. Two custom quality controls were prepared at levels that are on the sensitive part of the standard curve. Qualification criteria included as-sessment of the standard curve, quantification range, reproducibility (precision) and dilutional linearity (% recovery). Standard curve was made more robust by adding more points on the sensitive part of the curve. Lower limit of quantification was qualified to be statistically above the variance of the blank value. Reproducibility was good (%CV≤30%) among assays. Linearity was acceptable for kit standards diluted with castrated dog serum or commercially available serum matrix, also, for intact serum diluted with its respective castrated serum (R2>0.9). This assay can detect 8 fold INH-B difference between intact and castrated canine serum samples. Other parameters like frozen storage, freeze/thaw and lot-to-lot stability are pending. We conclude that this canine INH-B assay can consistently quantify INH-B levels in canine serum under the modified procedures.

1886 Mitigation of Fumonisin Biomarkers by Green Tea Polyphenols.
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Fumonisin B1 (FB1) is a carcinogen and a strong tumor promoter in animal models. Green tea polyphenols (GTP) are highly effective in inhibition of a variety of carcinogen-induced tumorigenic effects in many model systems. In this study we assessed mitigative effects of GTP on FB1-biomarkers in blood and urine samples collected from a randomized, double blinded, and placebo controlled intervention trial, which recruited a total of 124 people aged 20-55 who exposed FB1 via their corn-based diet. These participants were consented, randomly divided into 3 groups, and daily treated with either low-dose (GTP 500 mg, n=42), high-dose (1,000 mg, n=41) or placebo (n=41) for 3 months. Urinary levels of free FB1 at baseline were comparable (medium: 560.73, 574.56, and 559.09 pg/mg creatinine) for all three groups (p=0.162). Levels at urine samples collected at 1-month of the intervention was significantly decreased in the high-dose group (medium: 364.94 pg/mg creatinine; p<0.01) as compared with level in the placebo group (medium: 575.25 pg/mg creatinine). The inhibition rate is 18.95% in low-dose group and 33.62% in high-dose group. Levels of free FB1 at samples collected at 3-month of the intervention showed significant decrease in both low-dose (medium: 319.45 pg/mg creatinine; p<0.01) and the high-dose (medium: 215.85 pg/mg creatinine; p<0.01) groups as compared with the levels of the placebo group as well as the baseline levels. The inhibition rate is 40.18% in low-dose GTP group and 52.6% in high-dose GTP group. Levels of sphinganine (Sa), sphingosine (So), and their ratio in urine and serum samples were also evaluated in this study. These results demonstrate that supplement of GTP effectively mitigates urinary excretion of free FB1 via to be specified pathways in humans.

1887 Variation of Urinary Creatinine.
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Urinary creatinine has been commonly used for adjusting dilution status of urine species in biological monitoring. However, it can vary according to sex, age, race, BMI, meat intake, etc. The purposes of our study are to investigate the intra- and inter-individual variations of urinary creatinine in a sex, age and race matched subjects, and to study the impact of meat intake on the variations of urinary creatinine. We designed a diet-controlled study among the subjects who were Korean healthy females (N=9, age: 20±4 yrs, BMI 19.7±2.4 kg/m2) and measured urinary creatinine at 5 intervals during 24 hours with and without meat consumption. As results, diverse intra- and inter-variations of creatinine levels were shown in the subjects: When subjects did not take meat, the largest and smallest intra-variations in urinary creatinine ranges were detected in the subject C and G, i.e. 0.34-2.97 (Δ2.63)g/L and 0.93-1.63 (Δ0.7) g/L, respectively. With intake of meat (charcoal-grilled Korean beef tenderloin), the trend of intra-variation of urinary creatinine in each subject was not different (p=1.00 by Fisher exact test). It suggests that meat intake had little influence on intra- and inter-variation of urinary creatinine. In conclusion, our data re-emphasize that urinary creatinine must be measured in each spot urine even among the subjects who have similar age, sex, race and BMI due to its intra- and inter-variation. In the near future, the causes of intra- and inter-individual variations of urinary creatinine should be further studied.

1888 Cardiolipin As a Biomarker of Mitochondrial Dysfunction Associated with Parkinson's Disease.
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A commonly used pesticide, rotenone, is a mitochondrial respiratory complex I inhibitor. Rotenone accumulation was measured in brain homogenates of rotenone-poisoned C57BL/6 mouse brain tissues, and was qualified to be statistically above the variance of the blank value. Reproducibility was good (%CV≤30%) among assays. Linearity was acceptable for kit standards diluted with castrated dog serum or commercially available serum matrix, also, for intact serum diluted with its respective castrated serum (R2>0.9). This assay can detect >8 fold INH-B difference between intact and castrated canine serum samples. Other parameters like frozen storage, freeze/thaw and lot-to-lot stability are pending. We conclude that this canine INH-B assay can consistently quantify INH-B levels in canine serum under the modified procedures.
Parkinson disease (PD), we hypothesized that CL peroxidized molecular species ac- companying mitochondrial dysfunction/oxidative stress and apoptosis. In this study, we used circulating lymphocytes isolated from human blood and found that rotenone (50-250μM, 12-18h) caused apoptosis (phosphatidylserine external- 

ization, caspase 3/7 activation), reactive oxygen species production (superoxide, H2O2), mitochondrial dysfunction (inactivation of complex I, decrease of mito-

ochondria membrane potential, depletion of ATP) and activation of peroxidase ac-

tivity of mitochondria. Using an oxidative lipidomics approach, we found that treat-

ment of lymphocytes with rotenone resulted in accumulation of mono-lyso-CL and 

oxidized free fatty acids. In addition we were able to detect oxygenated mo-

lecular species of tetra-firinoe CL, a major CL molecular species in lymphocytes. 

Notably, molecular species of oxygenated CL formed in human lymphocytes were 
similar to those formed in cyt c driven reaction in the presence of H2O2 – in line 

with the known participation of cytochrome as a catalyst of CL peroxidation dur-

ing apoptosis. Using the combination of lipidomics and oxidative epitope-targeted 

enzymatic digestion of oxidized tetra-firinoe CL we found that its oxygenated LA 

species were represented by hydro- and hydroxy-derivatives. Thus, we con- 

clude that CL and its oxygenation products and metabolites may represent a new 
b biomarker of rotenone-induced mitochondrial dysfunction associated with PD. 

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1889 Evaluation of Insulin-Like Growth Factor Acid-Labile 
Subunit As a Novel Biomarker of Effect to the Mycotoxin 
Deoxynivalenol.

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Deoxynivalenol (DON) is a trichothecone mycotoxin produced from Fusarium 
species frequently found in grain products due to its recurrent contamination and 
resistance to food processing treatments. In growing experimental animals, chronic 
low-level DON exposure has resulted in anorexia, weight suppression and growth 
hormone axis perturbations. As a result, children are thought to be especially sensi-
tive to DON. Though a biomarker of exposure exists to measure DON exposure in 
humans, no biomarker of effect is currently available to predict the adverse negative 
weight effects of DON, thereby hindering complete risk assessment of this myco-
toxin. Two studies were conducted to assess the potential of plasma insulin-like 
growth factor acid-labile subunit (IGFALS) to be used as an effect biomarker for 
DON. In the first study, a 9 wk dietary DON exposure was employed in mice to 
test the hypothesis that depression in plasma IGFALS occurs at toxicologically rele-

vant doses prior to significant weight suppression. Results showed that the 1) NOAEL 
for depressed plasma IGFALS and weight was 2.5 ppm DON and 2) decrease 
plasma IGFALS was detectable before significant weight suppression was evident. 

In the second study, the specificity of reduced plasma IGFALS to DON, 
rather than DON-induced anorexia, was assessed using a dietary DON model. Mice 
were fed ad-lib control diet, restricted control diet or identical amounts of re-

stricted 15 ppm DON diet. Mice fed restricted DON diet exhibited significantly 
less plasma IGFALS than the restricted control indicating the specificity of plasma 
IGFALS reductions to DON. Thus, plasma IGFALS might be one suitable bio-

marker for predicting DON’s adverse growth effects in animals and humans.

1890 Validation of a Meso Scale Discovery Immunoassay for KIM-
1 Renal Biomarker in Cynomolgus Monkey (Macaca 
fuscularis) Urine.


The purpose of this study was to validate an immunoassay for detection of Kidney 
Injury Molecule 1 (KIM-1) in the urine of cynomolgus monkeys (NHP). 
Monkeys were treated with escalating doses of a compound and induced tubular 
degeneration/regeneration as determined by histopathology. Urine was collected 
predose, 16 days post dose and 21 days post dose and urine with low, medium and 
high levels of KIM-1 were used to validate the Meso Scale Discovery (MSD) 
Human KIM-1/TIM-1 (single-plex) immunoassay kit as this kit cross reacts with 
NHP KIM-1. Additional urinary samples from older colony monkeys as well as normal 
younger NHPs were also used to establish a preliminary observed range 
(<0.01ng/mL). We determined intra-assay (7.7% CV) and inter-assay precision 
(24.5% CV), limit of blank (0.00031ng/mL), limits of quantitation (0.01ng/mL to 
10ng/mL), dilutional linearity (not linear when diluted), recovery (84.6% - 122.8%), 
predictive quality control range evaluation (0.439ng/mL CV 18.7%, 0.615ng/mL CV 16.8%), freeze/thaw (F/T) stability out to 4 F/T cycles (95.5% - 121.5%), and sample storage stability out to 10 weeks. There was also a good bio-

logic correlation with time and dose-dependent increases in KIM-1 for toxicity 
study samples. All parameters measured showed acceptable immunoassay assay per-
f ormation and overall assay performance and the other results obtained, the validated method is robust and can be performed under good laboratory practice conditions to support nonclinical studies to assess for renal tox-
icity.

1891 Evaluation of a Three-Dimensional Oral Cell Model for the 
Assessment of Tobacco Products.

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Winston-Salem, NC. Sponsor: S. Theophilis.

Oral disease is frequently associated with viral and environmental exposures as well 
as oral hygiene. The goals of this study were to evaluate the impact of smokeless to-

bacco extracts (STE) and cigarette total particulate matter (TPM) on cell survival, 
oxidative stress, inflammatory response and tissue integrity using three-dimensional 
cultures of human buccal (EpiOral™) cells. 

EpiOral™ cells were treated with extracts of IS2 (reference dry snuff), 253 (refer-
ence moist snuff) and a smokeless tobacco blend prepared in complete artificial 
saliva (CAS) as well as with TPM from Kentucky Reference 3R4F cigarette 
(DMSO-based) for time points through 24 hours. Toxicity was assessed with the 
lactate dehydrogenase (LDH) assay. Glutathione (GSH) measurement and histo-

logical analyses were used to assess oxidative stress and changes in tissue integrity, 
respectively. Gene expression analyses were also conducted via qRT/PCR and mul-
tiples cytokine testing. 

Dose- and time-dependent release of LDH was observed for all test articles. The 
optimal exposure time appears to be 12 hours where 3R4F TPM elicited up to a 3-
fold increase in LDH release; the IS2 and 253 extracts yielded a 2-fold increase 
while no increase was observed for the smokeless tobacco blend. Tissue integrity 
was slightly disrupted by TPM exposure, while no impact was observed for the 
STE.

Oxidative stress as measured by GSH analysis was not apparent for any of the test 
articles; however, altered inflammatory response was observed by changes in IL-1α 
and G-CSF cytokine release and modulations in at least one of the following genes, 
IL-1α, TNFα or COX-2. The test articles also induced increases in cellular stress 
and oxidative stress, mitochondrial content and augmentation of caspase-3. 

Collectively, the data suggest that the EpiOral™ three-dimensional human cell cul-
ture model may be useful in evaluating tobacco extracts.

1892 Assessment of Cardiac Biomarkers in Cynomolgus 
Macaques.

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L. Tang, D. Fairchild, K. Lopez and J. C. Mirabela. SRI International, Menlo Park, 
CA.

A number of new cardiac biomarkers have recently been developed for use in ro-

dents; however, there are no validated cardiac biomarkers suitable for use in nonhu-

man primate (NHP) studies. We previously reported results of cardiac markers in 
African green monkeys (AGM) and Rhesus macaques (RM). In the current study 
we have extended these evaluations to Cynomolgus macaques (CM), the most 
widely used NHP species for toxicity studies. Two CM/sex were given a single sub-
cutaneous (sc) injection of isoproterenol (IPT; 4 mg/kg); 1 CM/sex received sc 
saline. Cardiac effects of IPT were observed within 1 hr postdose and included hy-
potension, ventricular premature complexes, ventricular bigeminy, atrial premature 
complexes, with or without aberrant conduction and ST segment elevation. Blood 
samples were collected predose and at 1, 4, 24, 48 and 72 hr postdose, and evalu-
ated with MSD MIP-1 muscle injury kits (rat: cTnI, cTnT, FABP3, Myl3, sTnI; 
human: TNI). IPT produced significant increases in the level of most cardiac bio-
markers: cTnI, FABP3, Myl3, sTnI and human cTnI were increased over predose 
levels by 4.2-, 2.5-, 25-, 28- and 23-fold, respectively, with peak times ranging from 4 to 8 hours. Similar results were seen in females, though rat cTnI was not 
increased in males, but a 4.9-fold increase was seen in females. At 72 hr postdose, 
there were still elevations in Myl3 and sTnI. IPT plasma levels at 1 hr postdose were 
higher in males (708 ng/ml) than in females (324 ng/ml) and fell to 324 and 302 ng/mL at 4 hr in males and females, respectively. Heart histopathology 3 days post-
dose revealed minimal to moderate cardiac myofiber degeneration, myofiber kary-

omegaly and leukocyte infiltration in all treated animals. These results indicate that 
the MSD rat and human muscle injury panel provides excellent sensitivity for as-
sessing cardiac effects in CMs, and these data are consistent with the utility of these 
kits previously reported in RM and AGM. Work supported by NIAID Contract 
N01-AI-70043.
Toxicogenomic approaches have identified protein biomarkers of renal cell injury/repair as early predictors of renal toxicity prior to changes in renal histopathology. We used these novel biomarkers to determine if rats orally dosed with industrial chelates exhibited altered urinary biomarker levels that preceded histopathologic changes in kidney. The nephrotoxicant/renal carcinogen, nitrilotriacetic acid (NTA), is known to cause rat proximal tubule cell injury/repair (3-7 weeks) followed by renal tumors (2 years) after oral dosing. A new, readily biodegradable chelate, L-glutamic acid diacetic acid (GLDA), previously showed no significant microscopic renal changes (90-day, oral), and EDTA, known non-carcinogen (oral bioassay) were also included in our study. Male Wistar rats were gavaged daily (oral, 28 days, 1000 mg/kg/day; n=10/group) with Na+ salts of these chelates. As expected, mean urinary levels of Na+ & Zn2+ were elevated in all chelate-treated groups. Two rats in NTA group were euthanized as moribund on Day 13. The surviving NTA group showed decreases in mean body weights, food and water consumption, and urine Mg2+, and increases in the mean levels of urine Ca2+, total protein, lactate dehydrogenase, kidney injury molecule-1 (Kim-1), Clutsterin (CLU) and increased proximal tubule cell proliferation (BrdU). No such changes were seen with GLDA. Kim-1 and CLU are inducible kidney proteins and are approved by FDA as predictive, early and noninvasive urinary biomarkers of kidney cell injury/repair. At necropsy, bilateral kidney enlargement (mean relative kidney weights) was noted with NTA, but not GLDA. In conclusion, our study showed that NTA, but not GLDA or EDTA, caused significant early renal cell toxicity when evaluated with urinary protein biomarkers as early predictors of nephrotoxicity.
1897 3, 3’, 4, 4’, 5-Pentachlorobiphenyl (PCB 126) Decreased the Ratios of Epoxide Metabolites to Their Corresponding Diols in Male Rodents.

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Oxylipins are oxygenated metabolites of certain fatty acid species. Changes in the homeostasis of these regulatory lipid mediators are of interest as markers of exposure to certain toxicants, including the aryl hydrocarbon receptor agonist 3, 3’, 4, 4’, 5-pentachlorobiphenyl (PCB 126). Here we test the hypothesis that chronic exposure to PCB 126 alters the levels of regulatory lipid mediators (oxylipins) in rats. Male Sprague-Dawley rats (5 weeks old) were treated biweekly with injections of PCB 126 in corn oil over a period of 3 months, representing cumulative PCB doses of 0, 0.06, 0.3, and 1.2 μg/kg body weight. PCB 126 treatment caused a dose-dependent reduction in growth and relative thymus weight, while relative liver weight was increased with PCB dose. PCB 126 levels in the liver increased in a dose-dependent manner, while PCB 126 levels in plasma were below the detection limit. The ratios of epoxide/diol metabolites displayed a dose-dependent decrease in plasma for oxylipins derived from arachidonic acid (ARA), eicosapentaenoic acid (EPA), docosahexaenoic acid (DHA) and linoleic acid (LA), as determined by metabolomics profiling of oxylipins with liquid chromatography-tandem mass spectrometry. Similarly, the epoxide/diol ratios for ARA and DHA derived oxylipins decreased in a dose-dependent manner in the liver. The effects of PCB treatment on epoxide/diol ratios were associated with significantly increased activities of soluble epoxide hydrolase (sEH) in cytosol and peroxisomes in the liver at the highest PCB 126 dose. Since increased sEH activity and decreased epoxide/diol metabolite ratios have been linked to cardiovascular disease and inflammation, our results suggest that changes in oxylipin plasma levels may be useful biomarkers of human exposure to PCB 126 and other dioxin-like compounds (supported by NIH grants ES04699 and E5013661).

1898 Identification of Neural Biomarkers of Altered Sexual Differentiation following Gestational Exposure.

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Sexual differentiation of the brain occurs during late gestation through the early postnatal period. The development of the phenotypical male brain is dependent on the aromatization of circulating testosterone to estradiol. Exposure to endocrine disrupting chemicals (EDCs) during early-life alters sexual maturation of the rat hypothalamic-pituitary-gonadal axis and subsequently, the timing of puberty and adult reproductive behavior. We set out to identify predictive neuronal biomarkers for use in evaluating the effect of EDCs on sexual differentiation in the male and female. We examined changes in gene expression in the rat hypothalamus [specifically, in the anteroventral periventricular nucleus (AVPV) and arcuate nucleus (ARC)] after in utero exposure to the aromatase inhibitor, letrozole. Pregnant dams were gavaged with vehicle or letrozole (0.1mg/kg) on gestational days 20 and 21. The AVPV and ARC were microdissected from male and female offspring on several pre-, peri-, and post-pubertal postnatal days (PND) and were evaluated for changes in expression of a number of neuropeptides that have sex specific patterns during development. These genes include those that encode for kispeptin (Kp) and tachykinin-2 (Tac2). We identified an increase in age with the expression of Kp in the AVPV of the genetic female, but not the genetic male. Kp expression in the AVPV of the letrozole-exposed genetic male was increased to the same level observed in the genetic female on PND 14, 25 and 49. However, Kp expression in the male ARC was not altered by letrozole. ARC Tac2 expression in the male was also not changed by letrozole. This work will provide optimal time points for these measures, but additional studies are needed to determine whether these and other CNS biomarkers, are predictive of altered puberty and/or sexual behavior due to early-life EDC exposure. This abstract does not necessarily reflect EPA policy.

1899 Biomarkers of Exposure and Effect in Long-Term Smokers and Moist Snuff Consumers.

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To assess the effects of chronic exposure to combustible and non-combustible tobacco product use, a single site, cross-sectional clinical study was conducted. Three cohorts of healthy males (40/cohort, 35-60 years) were enrolled: long-term smokers and moist snuff consumers (MSC), and non-tobacco consumers (NTC). Select biomarkers of exposure (BioExp) and potential biomarkers of effect (BioEff) indicating oxidative stress, inflammation and metabolic changes, among others, were investigated. Blood biomarkers were measured in subjects abstaining overnight from both food and tobacco, and urinary biomarkers were measured in ambulatory collections of 24h urine samples. Nicotine (and its metabolites) and total tobacco specific nitrosamines (TSNAs) were higher in MSC followed by smokers and NTC. Tobacco combustion-related BioExp (polycyclic aromatic hydrocarbons, aromatic amines and mercapturic acid metabolites) were significantly higher in smokers, with no difference observed between MSC and NTC cohorts. Compared to NTC and MSC, smokers exhibited significantly elevated levels of biomarkers associated with oxidative stress, inflammation, and platelet activation. Smokers also exhibited significantly higher levels of apolipoprotein B100 and oxidized low-density lipoprotein relative to NTC and MSC. Thus, alterations in BioEff suggesting inflammation, oxidative stress and altered lipid metabolism were detected in smokers compared to the non-smoking cohorts. In summary, our findings are in agreement with existing epidemiological data which show the reduced harm from smokeless tobacco consumption compared to smoking, with no-tobacco-use being the least risky. The BioExp and BioEff evaluated in this study are likely to be useful in future assessments of the health effects of new tobacco products.

1900 Mechanistic Biomarkers Provide Early Detection of Acetaminophen-Induced Acute Liver Injury at First Presentation.

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We investigated the potential of a panel of novel biomarkers - which demonstrate either enhanced liver expression or are linked to the mechanisms of toxicity - to identify patients with paracetamol (acetaminophen)-induced acute liver injury at first presentation to hospital when current liver injury markers are normal. In plasma samples from patients (n=129), at first presentation to hospital following paracetamol overdose, we measured miRNA-122 (mir-122, a liver specificity), High Mobility Group Box-1 (HMGB1; marker of necrosis), full length and caspase-cleaved Keratin-18 (K18; markers of necrosis and apoptosis, respectively) and glutamate dehydrogenase (GLDH; marker of mitochondrial dysfunction). In all patients, the biomarkers at first presentation significantly correlated with peak ALT/INR (International Normalized Ratio). In patients with normal ALT/INR at presentation, mir-122, HMGB1 and necrosis K18 identified the development of liver injury (n=15) or not (n=84) with a high degree of accuracy, and significantly outperformed ALT activity, INR and plasma paracetamol concentration for the prediction of subsequently liver injury (n=11) compared with no injury (n=52) in those patients presenting within 8 hours of overdose. Elevations in plasma mir-122, HMGB1, and necrosis keratin-18 identify subsequent development of acute liver injury in patients on admission to hospital, soon after paracetamol overdose, and in patients with ALTs in the normal range. The clinical development of such a biomarker panel could improve the speed of clinical decision making, both in the treatment of acute liver injury and in the design and execution of new treatment strategies that aim to refine the management of this common poisoning.

1901 Lead Concentrations Correlation between Fingernails and Whole Blood.

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Whole blood has been the biological fluid of choice to assess lead exposure. Other biomarkers need to be studied deeply, and the analysis of trace metals concentrations in nails is a non-invasive and simple collection procedure. The aim of this study was to investigate possible correlations between blood lead levels (BLL) and washed and non-washed fingernails lead levels (FLL) in 55 adults living in a lead contaminated area. Venous blood and fingernails (thumbs and forefingers) samples were collected. The nails of the left hand were washed with acetone and HCl 0.1 mol L-1 and the nails from right hand were not submitted to any pre-analytical procedure. Samples were analyzed by GF AAS and pairwise correlations were used to correlate lead concentrations between: BLL and FLL; nails from fingers from the
same hand and between washed and non-washed fingernails. It was found a non-significant correlation between BLL and washed thumb fingernail samples (r=0.219, p=0.112) and between BLL and thumb nails (r=0.182, p=0.191). Comparing fingernails from the same hand (thumb and forehead), lead concentrations of non-washed nails varied largely, even when transversal fragments from the same nail were analyzed. Lead concentrations in non-washed forefinger nails were not found to be correlated with washed thumb nails (r=0.169, p=0.189). On the other hand, for washed nails, thumb and forehead nails were found to be correlated (r=0.39, p=0.003). In conclusion, the results showed that non-washed nails are not a reliable biomarker for lead exposure. However, although washing up the nails may diminish the lead external contamination, the correlation between fingers is still weak to consider fingernail as a biomarker to lead exposure. In addition, even the washed nails were not found to be significantly correlated to BLL.

1902 Detecting and Quantifying Endogenous and Exogenous Formaldehyde Adducted Hemoglobin Utilizing Selected Reaction Monitoring.

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Widely recognized as a highly toxic compound, formaldehyde (FA) is considered to be a known animal and human carcinogen. It is ubiquitously present through normal cellular metabolism as well as from environmental pollutants. This compound is highly reactive towards macromolecules, forming diverse protein adducts and DNA damage which can be employed as biomarkers of chemical exposures. In particular, hemoglobin and red blood cells (RBCs) exposed to FA have been shown in laboratory studies to form an imidazolidone-type structure on the N-terminal peptides of hemoglobin alpha and beta chains. This data and the long lifetime of hemoglobin in RBCs (63 days in rats and 120 days in humans) afford investigation of the formation of endogenous and exogenous FA-hemoglobin adducts and the accumulation and loss of adducts. Monitoring this biomarker may reveal if inhaled FA enters the systemic circulation and reaches distant sites. In order to differentiate the origin of FA (endogenous versus exogenous), 10 ppm [13C12] FA rat inhalation exposures of 6 hours/day for 1 or 5 days were performed and globin was isolated from the washed and unwashed forefinger nails. Stable isotope labeled and unlabeled pairs of globin samples were synthesized and purified then reacted with FA to achieve the imidazolidone modification. These internal standards were then spiked into the globin samples prior to trypsinization and off-line fractionation. Utilizing protein cleavage isotope dilution mass spectrometry and selected reaction monitoring, methods were developed for quantification of the N-terminal valine FA-hemoglobin adducts and the accumulation and loss of adducts. Monitoring this biomarker may reveal if inhaled FA enters the systemic circulation and reaches distant sites.

1903 Assessment of the Relative Performance of Ten Urinary Biomarkers for Renal Safety across Twenty-Two Rat Toxicology Studies.


Novel kidney injury markers have been recently identified which may outperform or add value to the conventional kidney injury biomarkers blood urea nitrogen (BUN) and serum creatinine. To assess the relative performance of the growing list of these novel biomarkers, a comprehensive evaluation was conducted for 10 urinary biomarkers in 22 rat studies including both kidney toxicants and compounds with toxicities observed only in other organs. Furthermore, these kidney toxicity studies included proximal tubule toxicants as well as glomerular toxicants. The ten urinary biomarkers evaluated included Kim-1, Clusterin, Osteopontin, Osteocalcin, Albumin, Lipocandin-2, GST-alpha, β2-Microglobulin, Cystatin C, and Retinol Binding Protein 4 using novel immuno-based assays developed on the Mesoscale platform. Receiver operator characteristic (ROC) curves were employed as the main criteria to compare the performance of this panel of biomarkers in individual study animals against the microscopic histomorphologic changes observed. Of the kidney toxicity studies analyzed, Kim-1, Clusterin, and Albumin showed the highest overall sensitivity for detecting tubular injury, while Albumin exceeded all other markers in detecting glomerular injury. The data presented here represents a comprehensive parallel analysis of the performance of leading renal biomarker candidates, and further demonstrates that this multiplex approach enhances the ability to monitor drug-induced renal injuries as well as provide insight to linkages between individual biomarkers and specific histopathologic processes.

1904 Drug-Induced Kidney Injury Urinary Biomarker Response in Rats after Treatment with Nonnephrotoxicants.


Drug induced kidney injury (DIKI) biomarkers are important tools with which to monitor and diagnose acute and chronic kidney injury. Qualification of the biomarkers for use in nonclinical studies requires an understanding of the DIKI biomarker profile after treatment with nephrotoxicants and non-nephrotoxicants to fully understand the potential for false positives in future studies. Renally-acting pharmacotherapies were used in this study to investigate the renal biomarker profile of non-nephrotoxic drugs. Diuretic drugs were chosen to target specific nephron segments including furosemide (Loop of Henle), hydrochlorothiazide (distal tubule), spironolactone (collecting duct), and erythritol (pan-nephron). Male and female Han-Wistar rats (n=10) were treated orally for 14 days and biomarker excrion was measured on days 10, 16,22 (recovery n=5). Data were normalized to urine creatinine and then to control levels. There were no significant differences between controls and treated rats in excretion of any of the eight biomarkers assessed including: alpha-glutathione-s-transferase (αGST), mu-GST, renal papillary antigen-1 (RPA-1), clusterin, albumin, lipocandin-2, osteopontin and kidney injury molecule-1 (KIM-1). Osteopontin excretion in female rats and clusterin excretion in male rats were the most variable with up to 19.1 and 9.5 fold differences from controls, respectively, despite mean values being roughly equivalent to vehicle controls. The results from this study indicate that diuretic treatment affects different portions of the nephron does not result in increased biomarker signal. As such, the likelihood of obtaining false positive results due to physiological differences in animals in nonclinical safety screening studies is minimal.

1905 Urinary Biomarker Response to Hepsera-Induced Kidney Toxicity in Rats.

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Hepsera (adefovir dipivoxil) is an acyclic nucleoside phosphonate analog approved for treatment of hepatitis B infection. It is associated with renal tubular toxicity in rats and monkey, and has dose-limiting nephrotoxicity in the clinic. The purpose of this study was to evaluate urinary biomarker response for Hepsera-induced kidney toxicity in rats. Male Sprague-Dawley rats were first administered Hepsera at 20, 40 and 60 mg/kg/day orally for 14 days. Minimal to mild tubular degeneration was observed at 20 mg/kg/day with increased severity at 40 and 60 mg/kg/day. There were no changes in BUN or serum creatinine. In contrast, urinary biomarkers (KIM1, albumin, NAGL, and osteopontin) were dose-dependently elevated. To further investigate the time-course of the biomarker changes, male and female rats were treated with 20 mg/kg/day Hepsera for 1, 2, 4 weeks with 4 weeks of recovery. Tubular degeneration was observed with males more affected than females. This was first observed on Day 8 (minimal) and increased in incidence and severity with longer duration of dosing. Renal histopathology was still present at the end of recovery in males. There were no changes in BUN, while serum creatinine was slightly increased in males on Days 16 and 29. In males urinary KIM1, beta2-microglobulin (B2M), albumin, and NAGL were increased (2.5-37 fold) on Days 16 and 29. NAGL and B2M levels were still elevated after 29 days of recovery in contrast to KIM1 and albumin levels. Similar changes were observed in females on Day 29, but not on Days 8, 16 or recovery Day 29. These results indicate that urinary biomarkers can be more sensitive for Hepsera-induced kidney toxicity than BUN and serum creatinine.

1906 Novel Noninvasive Biomarker of Irradiation-Induced Gastro-Intestinal Injury.

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A non-invasive early marker of radiation induced gastro-intestinal (GI) injury continues to be in high demand. Hence, our aim was to develop a novel fecal biomarker of GI damage in mouse models which can be further extrapolated for use in non-human primates as well as in humans. A Reactive Nitrogen Species (RNS) detector compound NMAA-1 was established as a novel GI irritation/oxidative stress biomarker in an irradiated mouse model. NMAA-1 is a small molecule, which upon reaction with peroxynitrite (ONOO-) produces cleaved NMAA-1. The ability of NMAA-1 to detect ONOO- selectively and in a concentration-dependent manner in aqueous solution and in living cells is known. We explored this selectivity in the quantification of NMAA-1 and cleaved NMAA-1 in feces of mice with irradiation-induced GI damage. C57Bl/6 mice were irradiated at a high lethal dose of
13Gy or at a LD_{80/90} of 8.1Gy in two independent experiments using a Ca_{37} source. NMMA-1 was administered at 40-60 μM per mL via oral gavage without any signs of toxicity. Fecal samples were collected from the irradiated and non-irradiated control mice at different time points (pre-dose, 4, 6 and 8 hr post dose) prior to irradiation and also on days 1, 5 and 7 post irradiation from all animals. NMMA-1 and cleaved NMMA-1 levels were measured in the feces using LC-MS/MS method. Up to a three-fold increase in the level of the marker was noted in the fecal samples collected between days 5 to 7 in mice irradiated at 13Gy as compared to the control levels with C_{max} at about 6 hrs post dose. In case of 8.1Gy irradiation, two-fold or more increase in NMMA-1 and % cleaved NMMA-1 levels was seen in the feces until day 7. It was also observed that post irradiation the appearance of the fecal marker was moved to later time points probably due to decreased GI motility. NMMA-1 has proven to be very distinctive in its role as an indicative marker of GI damage due to radiation-induced toxicity. This promises to be an extremely important diagnostic tool for such condition and further optimization is ongoing.

1907 Exploring Challenges in Reconstructing Doses or Estimating Blood Concentrations from Urinary Biomarker Data Using a Pharmacokinetic Model for Perchlorate.


Many human biomarkers of current-use environmental chemicals are measured in urine. While urinary biomarkers are easier to collect than others (e.g., blood markers), reconstructing doses from urinary biomarkers is challenging. In many cases there exists a complicated relationship between dose and biomarker, for example, when a common biomarker for several chemicals exists and the concentration/relative proportion of the exposures from the environment are unknown. Even when there is a simple dose-biomarker relationship, many factors can impact the inter-predictability of urinary biomarker data, including urine volume, time between voids, and time between dose(s) and sample collection. In this study, a physiologically based pharmacokinetic (PBPK) model of perchlorate was used to examine how these various factors impact our ability to reconstruct doses from urinary biomarkers. The selection of perchlorate was based on its simple dose-biomarker relationship: 100% oral bioavailability and 100% excretion of the parent compound in urine. First, synthetic exposure profiles (varying daily intake doses, time of exposure, etc.) were generated and used as inputs to the PBPK model for simulating three biomarker sampling protocols: random spot voids, first voids, and 24 h cumulative voids. Correlations between simulated urinary concentrations and intake doses were calculated at both the individual- and population-level to determine which sampling protocol provides a more accurate and precise intake estimate. The strongest correlations were observed between the 24 h cumulative concentrations and average daily intake. Additional considerations for the impacts of changing urine volume, time between voids, and time between exposure and sample collection showed that random spot urine samples were highly sensitive to all three factors whereas 24 h cumulative samples were less sensitive.

[This abstract has been cleared by the EPA but solely expresses the view of the authors]

1908 Fecal and Blood Biomarkers for Gastrointestinal Injury and Subsequent Recovery.

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The search for reliable, non-invasive biomarkers of gastro-intestinal (GI) injury/recovery is ongoing but so far no biomarkers as reliable as histopathology have been identified in animal models. Our aim was to establish such biomarkers in non-human primates (NHPrs). Previously we established ELISA and MSD methods for lactoferrin and calprotectin measurements in NHPrs feces. These methods are currently being improved by multiplexing on one MSD platform. Two new markers including intestinal and liver fatty acid binding protein (I-FABP) and L-FABP were established as markers of intestinal epithelial integrity. These are released rapidly from GI enterocytes into the blood after cellular damage. I-FABP was assayed on a MSD platform for one species (specific antibody, antibody D), whereas L-FABP was analyzed using ELISA (LLOQ=102 pg/mL). Our MSD method for I-FABP has about 10 times lower LLOQ (9.7 pg/mL) as compared to its ELISA method (94 pg/mL). Normal ranges in NHPrs plasma were established and found to be above the LLOQ for both methods. These biomarkers proved to be helpful for assessment of drug therapeutic effect and potentially survival prognosis after irradiation GI damage. Additionally an LC-MS/MS method was established to estimate the systemic citrulline levels as a marker of GI injury. Citrulline is an amino acid released in circulation predominantly by the intestine enterocytes and is considered a marker of intestinal integrity. The LC-MS/MS method works in the dynamic range of 125-125000 ng/mL (~0.7-700 μM) of citrulline concentration in plasma. The L-FABP and I-FABP changes correlated well with the citrulline levels as well as histopathology findings based on the analysis of the samples obtained from a NHP irradiation study. Further optimization of the MSD, ELISA and LC-MS/MS methods are being undertaken to provide more sensitive assays from smaller sample volumes (including dried blood spot measurements) for these fecal and/or circulating biomarkers of gastro-intestinal inflammation, injury, and recovery.

1909 Clearance Rates of Organophosphate Metabolites in an Occupationally-Exposed Cohort of Farmworkers.

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Our studies in the Yakima valley of Washington state follow a cohort of 99 farmworkers (orchard workers) and 95 non-farmworkers to investigate potential exposures to organophosphate pesticides (OPs) by assaying urine samples for the 6 di-aryl non-specific metabolites of OPs. Urine samples were collected April-June 2005 on three dates spaced two days apart when OP pesticides were being applied and workers were in the orchards. The most commonly used OP in this period was azinphos-methyl and one of its metabolites, DMTP, had the highest concentration in urine compared to other metabolites of OPs for both farmworkers and non-farmworkers. We used a Bayesian Markov chain Monte Carlo method to estimate the joint distribution of the metabolites and clearance rates in a mixed effects model. The farmworker levels of DMTP were 19 times non-farmworkers. The clearance half-life of DMTP had a geometric mean (95% confidence interval) of 3.6 (2.5,6.0) days across all of the farmworkers (fixed effects), whereas non-farmworkers showed no clearance. The half-lives for individual farmworkers (random effects) varied between 1.5 days and no clearance. The shorter half-lives occurred in farmworkers who had higher first day concentrations of DMTP and farmworkers with no clearance with those who had lower first day concentrations. These results are consistent with a continuing variable level of occupational exposure of the farmworkers but some of the farmworkers having a higher exposure before our collection of urine. The pattern in the non-farmworkers is consistent with exposures to metabolites of OPs through diet and other routes of non-occupational exposure. Similar results were also found for DMP (Supported by grants P01 ES009601, P30 ES007033 from NIEHS and RD-834514, RD-831709, RD-832733 from US EPA. Contents are authors’ responsibility.)

1910 Assessment of MMP-3 and MMP-9 As Preclinical Biomarkers of Drug-Induced Vascular Injury.


Drug-induced vascular injury (DIVI) continues to be a major obstacle in drug development. Attempts to correlate this observation in preclinical toxicity studies with hemodynamic changes are not always successful. Additionally, no accessible and specific biomarkers of DIVI exist, making risk assessment and monitoring in humans a challenge during drug development and regulatory approval. It has been previously reported that circulating levels of matrix metalloproteinases (MMPs), specifically MMP-3 and MMP-9, were able to distinguish between patients with active antineutrophil cytoplasmic antibody (ANCA)-associated vasculitis (AAV) and remission. In order to determine whether MMPs are involved in the arterial pathology associated with DIVI observed in animal toxicology studies, a fluorescent imaging probe, MMPSenseTM, was used to measure activity of MMPs along with a fluorescent blood pooling agent, AngioSenseTM, ex-vivo in the mesenteric arteries of rats following administration of fenoldopam. Increased fluorescence in the mesenteric arteries from both imaging probes correlated well with vascular injury, as determined histologically, and was in agreement with increases in both MMP-3 and -9 expression in the affected arteries using immunohistochemical staining. To assess whether changes to circulating levels of MMP-3 and -9 respond to incidence and severity of DIVI, we compared their levels in plasma from several models of DIVI in rat, dog and monkey. Circulating levels of MMP-3 appear to correlate well with DIVI and the response is conserved across our pre-clinical species. In summary, MMPs appear to have a role in the arterial pathology associated with DIVI seen in animal toxicology studies, which is reflected in circulating levels of MMP-3 in pre-clinical species, and provide a potential accessible biomarker to monitor for DIVI in animals and in the clinic.
Molecular, Cellular, and Histological Changes in Mice Living on Sand Contaminated with Coal Dust under Laboratory Conditions.


Coal mining is one of the most important economical activities in Colombia. However, few data have been gathered concerning the impact of these activities on human and ecosystem health. During coal mining and transport, different types of dust material are formed. The aim of this work was to evaluate the toxic effects associated with the exposure to total suspended particle matter fraction of coal dust (<38 μm), from a sample collected in the Loma mine, Cesar, Colombia. Washed and sterilized sand was contaminated with coal dust to obtain concentrations ranging from zero (control) to 4%. Six different mice per group were allowed to live in boxes with this bed for eight weeks with ab libitum water and food. At the end of the experiment, animals were euthanized and blood and tissues collected. Mice in contact with coal dust did not evidence significant weight or hepatosomatic changes compared to control. However, animals in the 4% coal dust group grew faster. Real Time PCR analysis revealed an increase in Cyp1A1 mRNA for animals living on sand with concentrations greater than 2% coal dust. Unexpectedly, for mice on polluted sand, SOD and MTH hepatic mRNA were downregulated, and no changes were observed on Myc expression. The results of comet assay in peripheral blood cells, and the micronucleus test in blood smears revealed significant potential genotoxic damage at the greatest tested coal dust level. Histopathological analysis showed a dose-response relationship for the presence of hepatic necrosis, steatosis, vacuolization and nuclei enlargements in exposed animals. These results suggest that soil contaminated with coal dust induces several molecular, cellular and histopathological changes in mice. Accordingly, it is necessary to revise the current legislation on mining practices in Colombia in order to prevent health problems derived from these particles. Vice-Rectory for Research. UniCartagena. 2011-2012. Colciencias-UniCartagena, Colombia. Grant 110749326186 (2009).

Advancing Adoption of Novel Safety Biomarkers into Drug Development through Voluntary Submission of Data at US FDA, EMA, and PMDA.

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Regulatory science, the science of developing new tools, standards, and approaches to assess the safety, efficacy, quality, and performance of regulated medical products, has advanced over time due to a number of factors. However, many safety biomarkers used in nonclinical and clinical safety assessment in drug development have not changed in decades. Recently established channels for FDA, EMA, and PMDA to receive and evaluate scientific data supporting novel tools for use in drug development are now better defined in guidelines, e.g. FDA’s Draft Guidance for Industry “Drug Development Tool Qualification” and EMA’s “Evaluation of novel methods in drug development for use in drug development.” The outcome of a qualification submission to one of these regulatory agencies is a formal opinion regarding the utility of a novel drug development tool (DDT) for a particular and well-circumscribed application in the drug development process, defined within a “context of use.” While recognizing that other mechanisms exist within the research community for driving scientific consensus on novel biomarkers, this study focuses on the formal regulatory qualification process. This study compares the procedure, volume and types of submissions, and proposes a framework for assessing success at FDA, EMA, and PMDA. Safety biomarkers comprise over half of the sixteen unique biomarkers qualified thus far, and are highlighted. Qualification of new safety biomarkers by regulatory agencies and subsequent adoption by drug developers is anticipated to speed therapeutic development for patients in need, build scientific consensus as to the usefulness and readiness of novel tools for understanding disease and therapeutic development, and decrease uncertainty between the regulators and sponsors regarding their appropriate application.
Female Long Evans rats were maintained on these diets beginning 7 wk prior to breeding until the end of lactation. Dams were sacrificed on gestational day 16 or 20, or when pups were weaned on postnatal day (PN) 21. Fetuses were harvested with sacrifice of the dams, pups were sacrificed on PN14 and PN 21. Blood, thyroid gland, and brain were analyzed for iodine, TH, TH precursors and metabolites. Serum and thyroid gland iodine and TH were reduced in the two most deficient diets. TH was reduced in the fetal brain but was not altered in the neonate. Cognitive function, assessed by acoustic startle, water maze learning and fear conditioning, was unchanged in adult offspring, but excitatory synaptic transmission was impaired in the dentate gyrus by the two most deficient diets. A 15% reduction in cortical T4 in the fetal brain was sufficient to induce permanent reductions in synaptic function in the adult. These findings reflect potential of TH-disrupting chemicals, and suggest that standard behavioral assays do not readily detect neurotoxicity induced by modest developmental TH disruption. (Does not reflect EPA or CDC policy).

1916 Association of Paraoxonase-1 Activity and Gene Polymorphisms with Type 2 Diabetes Mellitus.

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Paraoxonase-1 (PON-1) is an HDL-associated lactonase that hydrolyses organophosphates such as paraoxon and diazoxon, the active metabolites of the pesticides parathion and diazinon, as well as affords protection against LDL oxidation. There is evidence that Type 2 diabetes mellitus (T2DM) patients display lower PON-1 activity. The objective of the present study was to determine the frequency of the PON-1192RR and 155QQ genotypes among patients and controls and to compare the serum levels of paraoxonase activity and paraoxonase-1 protein levels. We performed the genotyping with the polymerase chain reaction (PCR) method. We measured the paraoxonase activity in serum samples using the paraoxon-para-nitrophenol assay. There was no significant difference in the genotype distribution or in paraoxonase activity between the T2DM patients and healthy controls. The levels of paraoxonase activity found in serum were lower (p < 0.05) in T2DM patients compared to healthy controls, suggesting that PON-1 activity is reduced in patients with T2DM. The results of this study indicate that PON-1 activity and gene polymorphisms may be associated with the risk of developing Type 2 diabetes mellitus.


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Maternal exposure to 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) causes a number of developmental disorders including growth retardation in the pups. Although our previous studies have demonstrated that TCDD imprints defects in gender-specific phenotypes through a reduction in gonadotropin production in perinatal pups, the mechanism for growth retardation by TCDD remains unknown. In this study, we investigated a hypothesis that maternal exposure to TCDD damages maternal prolactin, a regulator of milk secretion and maternal care to the pups, and growth hormone (GH) in the pups to disturb the growth of offspring. Antihyperglycemic drugs have been reported to improve fetal growth retardation and GH levels. Therefore, our goal is to characterize the AhR knockout mouse (AhR-KO) and provide evidence for physiological roles for the AhR. We studied the physiological role of AhR in the liver, AhR knockdown (AhR-KO) mice with mice that express Cre recombinase under the control of the albumin promoter, resulting in loss of AhR expression specifically in the liver parenchyma. Our experiments indicate a novel sex dependent phenotype wherein, AhR-KO females exhibited reduced body weight, loss of adipose tissue and increased basal glucose levels (218 ± 19.73 mg/dl). Therefore, our goal is to characterize the AhR-KO mouse phenotype and identify potential AhR-dependent mechanisms responsible for lipodystrophy and abnormal glucose homeostasis. AhR-KO and control mice were subjected to glucose and insulin tolerance tests. We observed that AhR-KO females exhibited reduced body weight, loss of adipose tissue and increased basal glucose levels (218 ± 19.73 mg/dl). Therefore, our goal is to characterize the AhR-KO mouse phenotype and identify potential AhR-dependent mechanisms responsible for lipodystrophy and abnormal glucose homeostasis.

Perfluorooctane sulfonate (PFOS) is considered such as an endocrine disruptor. This study was designed to evaluate the possible alterations induced by PFOS on the hypothalamic-pituitary-testicular axis activity. For this purpose, male Sprague-Dawley rats were orally treated for 28 days with PFOS, at the doses of 0.5, 1.0, 3.0 and 6.0 mg/kg/day. Control rats received 0.5% Tween-20 vehicle. Twenty four hours after the last dose of PFOS, rats were killed by decapitation and the hypothalamus was removed in order to determine in this brain region two neuropeptide Y (NPY) and gonadotropin-releasing hormone (GnRH) concentration by specific enzyme-linked immunosorbent (ELISA) assays. Serum luteinizing hormone (LH), follicle stimulating hormone (FSH) and testosterone levels were measured by specific commercial kits. In addition, the relative gene expression of NPY and GnRH in hypothalamus of GnRH receptor in pituitary and of LH receptor and FSH receptor in testis was determined by quantitative real time PCR. Serum LH and testosterone levels and relative gene expression of GnRH and of FSH receptor were decreased in rats treated with PFOS. Serum FSH concentration and relative gene expression of LH receptor in testis were increased in these same animals. Hypothalamic concentration of NPY and GnRH was decreased with the dose of 1.0 and 3.0 mg/kg/day, but GnRH levels were increased with the dose of 6.0mg/Kg/day. Relative gene expression of NPY was diminished with the dose of 0.5mg/Kg/day, but it was increased with the dose of 6.0mg/Kg/day. Relative gene expression of GnRH receptor was not modified in pituitary by PFOS. The obtained results suggest that PFOS exposure can modify the hypothalamic-pituitary-testicular axis activity at several levels, and these alterations seem to be dependent of the administered dose. This work was supported by a grant from the Ministry of Education and Science, Spain (AGL2009-09061).

1919 Conditional Knockout of the Argyro Hydrocarbon Receptor in the Liver Alters Mouse Phenotype As Well As Glucose and Lipid Homeostasis.

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The aryl hydrocarbon receptor (AhR) is a ligand-activated transcription factor historically known for its role in the adaptive metabolism of xenobiotics. However, these developmental disorders including growth retardation in the pups, and growth hormone (GH) in the pups to disturb the growth of offspring. Antihyperglycemic drugs have been reported to improve fetal growth retardation and GH levels. Therefore, our goal is to characterize the AhR knockout mouse (AhR-KO) and provide evidence for physiological roles for the AhR. We studied the physiological role of AhR in the liver, AhR knockdown (AhR-KO) mice with mice that express Cre recombinase under the control of the albumin promoter, resulting in loss of AhR expression specifically in the liver parenchyma. Our experiments indicate a novel sex dependent phenotype wherein, AhR-KO females exhibited reduced body weight, loss of adipose tissue and increased basal glucose levels (218 ± 19.73 mg/dl). Therefore, our goal is to characterize the AhR-KO mouse phenotype and identify potential AhR-dependent mechanisms responsible for lipodystrophy and abnormal glucose homeostasis. AhR-KO and control mice were subjected to glucose and insulin tolerance tests. We observed that AhR-KO females exhibited reduced body weight, loss of adipose tissue and increased basal glucose levels (218 ± 19.73 mg/dl). Therefore, our goal is to characterize the AhR-KO mouse phenotype and identify potential AhR-dependent mechanisms responsible for lipodystrophy and abnormal glucose homeostasis.
Research on endocrine modulatory effects of Cadmium (Cd) started over a decade ago, when this metalloestrogen was found to interact with the estrogen signaling pathway. Since then, several independent in vitro as well as in vivo reports have emerged on this topic. Our objective has been to characterize the details of molecular mechanisms of action for the endocrine modulatory effects of Cd. We applied a combined in vivo and in vitro approach by using transgenicERE-luc reporter mice model as well as hepG2, MCF-7 and ECC-1 cell lines. After 3-days s.c. exposure to CdCl₂, we did not detect reporter gene activation in the dose ranges 5-500 μg/kg bw and 0.5-500 μg/kg bw in female and male mice respectively. Nevertheless, we observed significant thickening of the uterine epithelium in the absence of uterine weight increase in females, and detected activation of Raf, Erk1/2 MAPK in the liver of both genders in low dose groups. Further, in our in vitro settings, low doses of CdCl₂ (1nM-100nM) also activated Raf, Erk1/2 MAPK and this effect disappeared with the inhibition of GPR30 and EGFR receptors. Our data shows that the molecular markers that are modulated by Cd differ depending on the exposure level; i.e. low doses relevant to human exposure via diet stimulate cytoplasmic kinases, while higher doses induce cellular stress responses. We conclude that CdCl₂ affects cellular signaling pathways that can produce physiological effects reminiscent of bonafide estrogen stimulation. However, CdCl₂ does not activate canonical estrogen signaling.

**1921 Bisphenol A (BPA) Levels Were Associated with Increased Estrogen.**

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**BACKGROUND:** The plastic associated compound bisphenol A (BPA) is a known estrogen-receptor agonist. Since background exposure to BPA could be detected in most individuals, we explored the relationships between BPA levels in serum and levels of endogenous sex hormones and their precursors in a population-based sample.

**METHODS:** 1,016 subjects all aged 70 years were investigated in the Prospective Investigation of the Vasculature in Uppsala Seniors (PIVUS) study. BPA was measured in serum at ALS Canada using an API 4000 liquid chromatography/tandem mass spectrometry system. Serum samples were assayed by high-resolution chromatography coupled to tandem mass spectrometry. This study evaluates the concordance of serum and urine steroid concentrations as quantified by both RIA and LC-MS/MS following exposure to a known EDC, atrazine (ATR). Adult male rats were dosed with ATR (200 mg/kg/d) or methylcellulose (solvent control) by oral gavage for 5 days. Animals were decapitated 2 hours after the final dose. Serum was collected and separated into 2 aliquots for analysis. Serum was assayed by RIA for androstenedione (A), corticosterone (CORT), estradiol (E2), estrone (E1), progesterone (P4), and testosterone (T). Serum was extracted via solid phase extraction prior to LC-MS/MS analysis with positive electrospray ionization in multiple-reaction monitoring mode for A, CORT, P4, and T. E1 and E2 serum concentrations were quantified similarly by LC-MS/MS, following derivatization with danyl chloride. To compare CORT values from urine, pregnant adult rats were dosed with either ATR (100 mg/kg/d) or methylcellulose by oral gavage for 5 days (i.e., gestational days 14-18). Urine samples were collected daily for 2 consecutive 6 hour intervals following dosing and assayed for CORT by RIA and LC-MS/MS as described above. Data analyses demonstrated a high degree of correlation between the two detection methods (R2 = 0.88 – 0.92). No statistically significant differences were observed between RIA and LC-MS/MS means for any of the steroids assayed. These findings indicate that steroids may be reliably measured in rat biological media using RIA or LC-MS/MS in toxicological studies.

**PS 1924 P-p-DDE Levels Are Associated with Reduced Testosterone Levels in Elderly Males.**

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**Background:** The DDT metabolite p-p-DDE is a known androgen-receptor agonist. Since background exposure to p-p-DDE still could be detected in most individuals, we explored the relationships between p-p-DDE levels in serum and levels of endogenous sex hormones and their precursors in a population-based sample. Methods: 1,016 subjects all aged 70 years were investigated in the Prospective Investigation of the Vasculature in Uppsala Seniors (PIVUS) study. p-p-DDE was measured in plasma by a high-resolution chromatography coupled to tandem mass spectrometry (HRGC/HRMS), and sex hormones by high-specificity liquid chromatography/tandem mass spectrometry. Results: In men, p-p-DDE levels were related to both reduced testosterone (p=0.006) and SHBG levels (p=0.008). Furthermore, also reduced levels of the precursors pregnenolone and 17-OH-pregnenolone were related to high p-p-DDE levels (p=0.05), indicating a general reduction in endogenous sex hormone synthesis in subjects with high p-p-DDE levels. Similar trends, but less pronounced, were seen in women.

**1922 A Comparison of RIA and LC-MS/MS Methods to Quantify Steroids in Rat Serum and Urine following Exposure to an Endocrine Disrupting Chemical.**

B. W. Riffle1,2, S. C. Lucas1 and W. M. Henderson3

Commercially available radioimmunoassays (RIA) are frequently used in toxicological studies to evaluate effects of endocrine disrupting chemicals (EDCs) on steroidogenesis in rats. Clearly there is a discrepancy comparing steroid concentrations in rats as measured by RIAs to those obtained using liquid chromatography coupled with tandem mass spectrometry (LC-MS/MS). This study evaluates the concordance of serum and urine steroid concentrations as quantified by both RIA and LC-MS/MS following exposure to a known EDC, atrazine (ATR). Adult male rats were dosed with ATR (200 mg/kg/d) or methylcellulose (solvent control) by oral gavage for 5 days. Animals were decapitated 2 hours after the final dose. Serum was collected and separated into 2 aliquots for analysis. Serum was assayed by RIA for androstenedione (A), corticosterone (CORT), estradiol (E2), estrone (E1), progesterone (P4), and testosterone (T). Serum was extracted via solid phase extraction prior to LC-MS/MS analysis with positive electrospray ionization in multiple-reaction monitoring mode for A, CORT, P4, and T. E1 and E2 serum concentrations were quantified similarly by LC-MS/MS, following derivatization with danyl chloride. To compare CORT values from urine, pregnant adult rats were dosed with either ATR (100 mg/kg/d) or methylcellulose by oral gavage for 5 days (i.e., gestational days 14-18). Urine samples were collected daily for 2 consecutive 6 hour intervals following dosing and assayed for CORT by RIA and LC-MS/MS as described above. Data analyses demonstrated a high degree of correlation between the two detection methods (R2 = 0.88 – 0.92). No statistically significant differences were observed between RIA and LC-MS/MS means for any of the steroids assayed. These findings indicate that steroids may be reliably measured in rat biological media using RIA or LC-MS/MS in toxicological studies.

This abstract does not reflect U.S. EPA policy.
In conclusion, increased levels of p-p-DDE levels were associated with both reduced testosterone and SHBG levels in elderly males, indicating endocrine disrupting activity by DDT in the elderly.

Furthermore, there are often knowledge gaps in the studies used to assess EDCs. To address these concerns, we performed a pre-/post-natal reproductive toxicity study of vinclozolin as part of a larger BASF and CEFIC funded project to measure the developmental toxicity of low single- and mixture-doses of anti-androgens. The tested doses were selected to mimic a low-effect level, the no observed adverse effect level (NOAEL) for endocrine effects, and the acceptable daily intake (ADI). At the LOAEL dose, female offspring developed normally, while the male offspring showed effects known for anti-androgens. No effects at all were noted at the lowest two doses, the NOAEL and the ADI. At the top dose, an increase was noted in nipples and/or areolas in male animals on PND 12. This effect was transient, as all had regressed by PND 21. In both of these dose groups, male offspring which were reared to young adulthood displayed additional anti-androgen effects including delayed sexual maturation and reduced male sex organ sizes and weights. However, no significant decrease in ano-genital distance on PND 1 was noted in the animals of this dose group. Similarly, assessment of sexual steroid hormones and their precursors revealed no effects at any of the dose levels. Taken together, the weight of evidence of the clinical and pathological findings suggests a lack of a non-monotonic dose-response curve.

Endocrine disruption has become an important topic of public concern. Despite an increasing amount of attention, little is understood about if environmentally relevant doses of endocrine disrupting chemicals (EDCs) affect homeostasis.

In summary, we performed a pre-/post-natal reproductive toxicity study of vinclozolin as part of a larger BASF and CEFIC funded project to measure the developmental toxicity of low single- and mixture-doses of anti-androgens. The tested doses were selected to mimic a low-effect level, the no observed adverse effect level (NOAEL) for endocrine effects, and the acceptable daily intake (ADI). At the LOAEL dose, female offspring developed normally, while the male offspring showed effects known for anti-androgens. No effects at all were noted at the lowest two doses, the NOAEL and the ADI. At the top dose, an increase was noted in nipples and/or areolas in male animals on PND 12. This effect was transient, as all had regressed by PND 21. In both of these dose groups, male offspring which were reared to young adulthood displayed additional anti-androgen effects including delayed sexual maturation and reduced male sex organ sizes and weights. However, no significant decrease in ano-genital distance on PND 1 was noted in the animals of this dose group. Similarly, assessment of sexual steroid hormones and their precursors revealed no effects at any of the dose levels. Taken together, the weight of evidence of the clinical and pathological findings suggests a lack of a non-monotonic dose-response curve.

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**1930 BPA and TBBPA Target ATP Binding Cassette Transporters in the Blood-Testis Barrier and Impair Leydig Cell Steroidogenesis.**

M. J. Roelofs1, A. C. Dankers2, A. H. Piersma1, F. G. Russel1, R. Masereeuw1 and M. van Duursen3

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Disturbances of androgen production in Leydig cells and/or androgen function in Sertoli cells in the (female) testis have been shown to cause a wide variety of male reproductive abnormalities in experimental animals. Drug transporters in the blood-testis barrier (BTB) prevent entry and accumulation of xenobiotics in the testis, but are also involved in local transport of steroid hormones. Among these are the ATP binding cassette (ABC) transporters, P-glycoprotein (P-gp/ABCB1), breast cancer resistance protein (BCRP/ABCG2), multidrug resistance protein 1 (MRP1/ABCC1) and 4 (MRP4/ABCC4). We studied the interaction of Bisphenol A (BPA) and Tributylmethoxobisphenol A (TBBPA) with human P-gp, BCRP MRP1 and MRP4-mediated transport using membrane vesicles of human embryonic kidney (HEK293) cells overtaking these transporters. Also, the effects of these compounds on testosterone production and expression of steroidogenic genes (Star, Cyp11a1, 3β-HSD, 17β-HSD, Ins3, Cyp17A, Cyp19, LH receptor, Sd5a1) in mouse Leydig MA-10 cells were determined. BPA and TBBPA concentration-dependently induced testosterone secretion by MA-10 cells and expression of specific mRNAs of 17β-HSD and Sd5a1 (5α-reductase case 1) were upregulated. BPA and TBBPA differentially inhibited P-gp, BCRP MRP1 and/or MRP4 activity. Blocking P-gp using PSC833, increased testosterone secretion upon BPA and TBBPA exposure. The MRP inhibitor MK771 completely blocked testosterone secretion elicited by BPA and TBBPA. Our data suggest that EDCs might disrupt local androgen production and androgen transport from the Leydig into Sertoli cells thus affecting normal germ cell development.

**1931 Effects of GPER Activation on (Xeno) Estrogen-Induced Cellular Responses.**

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Estrogen can exert cellular effects through both nuclear (ERα and ERβ) and membrane-bound receptors (GPER). It is unclear if these receptors act independently or engage in crosstalk to influence hormonal responses. To investigate each receptor’s role in proliferation, activation of reporter genes and protein phosphorylation in breast cancer cells (MCF-7), we employed ESR1 (PPT), ESR2 (DPN) and GPER (G1) selective agonists. As anticipated, 17β-estradiol (E2), PPT and DPN, enhanced cell proliferation, whereas G1 had no impact. However, E2, PPT- and DPN-induced proliferation was repressed when cells were co-exposed with G1. Similar results were observed for activation of an estrogen response element (ERE)-driven luciferase reporter gene where G1 elicited a dose-dependent decrease in E2-, PPT- and DPN-induced activity. As membrane receptors typically initiate a series of phosphorylation events, we performed global cytosolic phosphoproteome analysis driven luciferase reporter gene where G1 elicited a dose-dependent decrease in E2-, PPT- and DPN-induced activity. As membrane receptors typically initiate a series of phosphorylation events, we performed global cytosolic phosphoproteome analysis. Of the 238 phosphorylated proteins identified only 10 sites were shared between E2 and G1 whereas 31 and 13 phosphorylation events were specific for each ligand, respectively. Using pathway analysis we developed a putative network of proteins that are influenced by GPER activation and have identified ‘central nodes’ (i.e. NDRG2) which highlights a possible role for these proteins as targets of GPER activity. As xenoestrogens, such as bisphenol-A and genistein, bind to ESRs and GPER we are currently investigating the impact these compounds have on nuclear and membrane estrogen receptor signaling networks. Overall, our proteomic and mechanistic approaches will lead to the identification and sorting of GPER network and the potential interplay with E2R that will illuminate the role of these receptors in cellular responses to E2 and xenoestrogens.

**1932 Antiproliferative, Antiandrogenic, and Cytotoxic Effects of Novel Synthetic Derivatives of Caffeic Acid in LNCaP Human Androgen-Dependent Prostate Cancer Cells.**

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Caffeic acid and its naturally occurring derivative caffeic acid phenethyl ester (CAPE) have antiproliferative and cytotoxic properties in a variety of cancer cell lines, but little is known about their effects on prostate cancer cells. We evaluated the effects of caffeic acid, CAPE, and 18 novel synthetic derivatives on cell proliferation, subcellular localisation and expression of androgen receptor (AR) and secretion of prostate-specific antigen (PSA) in LNCaP human hormone-dependent prostate cancer cells. LNCaP cells cultured in steroid-free medium were exposed to 0.1 nM dihydrotestosterone (DHT) in combination with various concentrations of caffeic acid derivatives (0.3-100 μM) for 24-72 h during which cell proliferation was followed using an xCELLigence cell monitoring system (Roche). Cytoplasmic and nuclear levels of AR were determined by immunoblotting and PSA secretion using a commercial ELISA. Cytotoxicity was measured by assessing mitochondrial function using a WST-1 assay. Seven synthetic derivatives of CAPE were strong, concentration-dependent inhibitors of androgen-stimulated LNCaP cell proliferation with up to 3-fold greater potencies (IC50=8.24-2.4 μM) than CAPE (IC50=28.29±4.5 μM): caffeic acid had no effect. Concomitant with inhibition of cell proliferation, DHT-stimulated PSA secretion was reduced by CAPE and the 7 derivatives. The most potent derivative MT-30 (IC50=7.9 ± 2.4 μM) inhibited DHT-stimulated cell proliferation and PSA secretion significantly at 0.3 μM. Exposure to 10 nM DHT increased cytoplasmic and nuclear AR levels and co-treatment with increasing concentrations of MT-30 and CAPE, interestingly, further increased these levels. In conclusion, a number of synthetic derivatives of caffeic acid are potent inhibitors of androgen-dependent prostate cancer cell growth, acting via an antiandrogenic mechanism that involves increased nuclear accumulation of (possibly inactive) AR.

**1933 Cadmium Content in Human Pancreatic Insulin Producing Beta Cells.**

M. El Muayed1, W. W. MacRenaris2 and W. L. Lowe1

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Evidence suggests that chronic low level cadmium exposure impairs the function of insulin producing beta cells and may be associated with type 2 diabetes mellitus. We previously reported that insulin producing beta cells accumulate Cd avidly and gradually in a dose and time dependent manner over prolonged time periods, leading to impaired beta cell function at environmental Cd exposure concentrations. This is likely due to the high beta cell Zn turnover and the significant overlap between Zn and Cd transporters in these cells. In the present study we compared the content of cadmium (Cd) and Mercury (Hg) between native human islets from type 2 diabetes patients with that of liver samples (10 human subjects living in the USA). The content of Cd and Hg in these tissues lysates was measured by ICP-MS. All studies were approved by our institutional review board. The cadmium (Cd) content in islets from 10 human non-diabetic subjects ranged from 7.4 to 71.92 nM/g protein (average 28.80 ± 6.73 nM/g protein). The concentration of mercury (Hg) was significantly lower ranging from undetectable (detection threshold 4 nM/g protein) to 5.05 nM/g protein. In comparison, human liver lysates contained significantly less Cd (9.04 ± 1.39 nM/g protein). No Hg was detected in any of the liver samples (detection threshold 4 nM/g protein). We conclude that human islets accumulate measurable, and likely relevant quantities of Cd that are higher than these of Hg under normal environmental exposure conditions. In comparison, the Cd concentration in Liver samples was significantly lower, likely due to the significantly lower Zn turnover rate in liver cells compared to that in beta cells.

**1934 Formulation Development and Analysis Methods for Bisphenol AF in 5K96 Verified Casein Feed.**

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Bisphenol AF (BPAF), a perfluorinated BPA homologue, is a cross linking agent in fluoroelastomers used in food and pharmaceutical processing equipment. Due to inadequate toxicity data, occupational and potential consumer exposure, and structural similarity to bisphenol A, BPAF was selected for evaluation by the National Toxicology Program (NTP). MRGlobal supported the NTP by formulating BPAF
into a phytoestrogen-free rodent diet, LabDiet 5K96 Verified Casein diet, developing the analysis of the effects on metabolic activity, and evaluating homogeneity and storage stability in the formulated diet.

The analysis method involved the extraction of BPAF from feed using acetonitrile-acetic acid (99/1 v/v). The method covered a formulation range of -200 to -10,000 μg/g with dilution into the curve up to 15,000 μg/g. The method was linear, accurate, and precise with BPAF recoveries ≥ 99%. Homogeneity evaluation of 97.5 and 15,000 μg/g formulations showed recoveries ≥ 99% with RSD values ≤ 0.9%. Long-term BPAF stability for the 93.5 μg/g formulation showed the formulation was stable for 42 days (recovery ≥ 94.5%) when stored under refrigerated or freezer conditions. When the 93.5 μg/g formulation was stored under simulated dosing conditions, in the presence of rat urine and feces, the ACN-acetic acid extraction resulted in an 68% BPAF recovery after 7 days, indicating possible BPAF instability or extensive binding to feed under these conditions. To investigate this, several solvents with different polarity and pH were tested. ACN/acetic acid 99/1 extraction resulted in 79% BPAF recovery after 8 days storage in the presence of rat urine and feces. Adding an additional acid-digestion and extraction step, in which the formulation was digested with 3.8N HCl at ~ 75°C and extracted with ethyl ether:petroleum ether (1:1), BPAF recovery increased to 90% in Day 7 samples. This suggests that BPAF is stable but exhibits non-covalent binding to feed components.

**1935 Silencing of Testisin through CpG Methylation from Exposure to Phthalates in Ntera-2 Cells.**

J. Gomes, J. Kapingo, B. Nguyen and D. Krewski. University of Ottawa, Ottawa, ON, Canada.

Di (2-ethylhexyl) phthalate (DEHP) is one of the most highly produced and frequently studied phthalates. Its metabolite, monoester mono (2-ethylhexyl) phthalate (MEHP) is reported to be a testicular toxicant. Following toxicity pathway analyses we identified Testisin, GSTP1 and MGMT genes to study their expression in testicular germ cells (Ntera-2). Testisin present in normal tissue but absent in neoplastic tissue and regulates proteolytic reactions in testicular germ cells; GSTP1 inactivates carcinogens and is a member of glutathione-S-transferase; and Methylguanine DNA methyltransferase (MGMT) provides defense against neo- plasm. Testicular cells under laboratory conditions were exposed to MEHP in a dose- and time-dependent manner at concentrations of 1μM, 10μM, and 100μM at 24, 48, 72 and 96th time points. The control was made by exposure of MEHP to 5’aza (demethylating agent) for hypermethylation. After exposure, those genes were analyzed by Quantitative Real Time PCR (qRT-PCR). The results revealed an overall down regulation for each gene as the concentration and/or time increased. The expression of Testisin, GSTP1 and MGMT was downregulated but not significantly in the last two cases. Following exposure to 5’aza treatment and coexposure, there was a significant up-regulation and restoration of the expression of the Testisin gene. This suggests that MEHP may down-regulate Testisin gene expression by DNA methylation. These findings provide evidence that MEHP can alter the expression of Testisin, GSTP1 and MGMT, genes. Testisin which is associated with testicular germ cell tumors and its downregulation and subsequent restoration may be caused by DNA methylation following exposure to MEHP. The expression of GSTP1 which is a xenobiotic metabolizing enzyme gene and the DNA repair enzyme gene suggests that the toxicant if fairly active in these cells exposed to MEHP. The investigation of DNA methylation at the CpG islands of the promoter region of Testisin is described.

**1936 An Examination of the Effects of Methyltriclosan on Early-Embryonic Development in the South African Clawed Frog (Xenopus laevis).**

M. Crombie1, M. Wages2, B. Perafan1, E. Smith2 and J. Carr1. 1Biological Sciences, Texas Tech University, Lubbock, TX; 2The Institute of Environmental and Human Health, Texas Tech University, Lubbock, TX. Sponsor: W. Guo.

Triclosan (5-chloro-2-(2,4-dichlorophenoxy)phenol) is a commonly used bactericide present in many personal care products such as detergents, liquid hand soaps, deodorants, cosmetics, lotions, mouthwash, and toothpaste, and it can be integrated with fabrics, plastics, carpets, and toys. Methyltriclosan is a derivative that is formed from triclosan via biological methylation at an unknown interval during waste water treatment. Methyltriclosan is more abundant in the environment, more lipophilic than triclosan, and has a greater potential to accumulate in fatty tissues. The global decline of amphibian populations has raised awareness surrounding the possible effects of poor water quality on the health of habitats. Since metamorphosis and reproductive development in amphibians is highly regulated by thyroid hormone (TH), and the structure of triclosan is similar to that of TH, there is the possibility that triclosan and methyltriclosan may act on TH receptors to alter metamorphosis and reproductive development. Standard 96-hour Frog Embryo Teratogenic Assay-Xenopus protocols were followed using South African Clawed Frog (Xenopus laevis) embryos. After measuring the larvae the data revealed that early embryonic exposure to environmentally relevant concentrations of methyltriclosan resulted in statistically significant alterations in total body length, snout-to-vent length and crown width. Furthermore, molecular studies were performed to identify the effects of methyltriclosan exposure on the TH-responsive gene, THßZIP. The results of the quantitative real-time polymerase chain reaction did not support the induction of THßZIP gene expression after exposure to environmentally relevant concentrations of methyltriclosan. However, the expression of other TH-responsive genes may be altered upon exposure. Collectively, these data are the first to report on the responsiveness of vertebrate embryos to methyltriclosan exposure.

**1937 No Evidence of Endocrine Disruption by Glyphosate in Male and Female Pubertal Assays.**

J. Bailey1, 2, J. Hauswirth3 and D. Stump1. 1 Dow AgroSciences, LLC, Indianapolis, IN; 2Joint Glyphosate Task Force, Raleigh, NC; 3WIL Research Laboratories, LLC, Ashland, OH.

The Food Quality Protection Act and Safe Drinking Water Act amendments (1996) required the USEPA to develop the Endocrine Disruptor Screening Program (EDSP), which currently consists of 11 Tier 1 screening assays to evaluate the potential for a chemical to interact with the endocrine system. Glyphosate was included in the first list of 67 compounds subject to the EDSP, which were selected for screening based on their potential for exposure rather than suspected interference with the endocrine system. The potential for glyphosate (G) to induce endocrine disruption has now been evaluated in the male and female Pubertal assays, the two most apical mammalian EDSP Tier 1 assays. The female pubertal assay evaluates potential effects on pubertal development and thyroid function in the juvenile female. Four groups of fifteen juvenile female rats were dosed with the following: 0, 100, 300 and 1000 mg/kg G once daily via oral gavage from postnatal day (PND) 22 to 42. There was no evidence of any direct test substance-related estrogenic or anti-estrogenic effects, nor was there any evidence of direct test substance-related effects on pituitary development or thyroid function in the juvenile female rat following oral administration of glyphosate at any dosage level tested. The male pubertal assay evaluates potential effects on pubertal development and thyroid function in the juvenile males. Four groups of fifteen juvenile male rats were dosed with the following: 0, 100, 300 and 1000 mg/kg G once daily via oral gavage from PND 23 to 53. There was no evidence of any direct test substance-related androgenic or anti-androgenic effects, nor was there any evidence of direct test substance-related effects on pubertal development or thyroid function in the juvenile male rat at any dosage level tested.

Based on these results, glyphosate does not exhibit endocrine disruption in the male and female pubertal assays.

**1938 Exposure to G-1, a Selective Agonist for G Protein-Coupled Estrogen Receptor 1 (GPER), Results in Elevated Levels of Vitellogenin in Adult Fathead Minnows (Pimephales promelas).**

S. Jayasinghe1, K. Kroll1, N. Denslow1 and T. Sabo-Attwood1. 1Department of Physiological Sciences, Center for Environmental and Human Toxicology, University of Florida, Gainesville, FL; 2Department of Environmental and Global Health, Center for Environmental and Human Toxicology, University of Florida, Gainesville, FL.

Several research groups have shown that G protein-coupled estrogen receptor-1 (GPER) mediates 17β-estradiol (E2) activation through non-genomic membrane initiated pathways. Estrogens play critical roles in a variety of biological processes, including reproduction and development in vertebrates. In fish, hepatic synthesis of vitellogenin (VTG), a precursor egg yolk protein that is vital to successful reproduction, is controlled by E2 via nuclear estrogen receptors (ESR). However, the involvement of GPER in vitellogenesis has not been investigated. As the GPER is not a well characterized in fish the aims of our study are to: (1) assess the tissue-specific expression of GPER in adult fathead minnows (FHM) and (2) determine the effect of GPER activation on VTG synthesis in FHM males and females using a selective GPER agonist (G-1) and antagonist (G-15). Using qRT-PCR we show that GPER mRNA is detectable in numerous organs, and is most highly expressed in the brain followed by gall bladder, trunk kidney, intestine, liver, heart, ovary and muscles. Aqueous exposure to G-1 (5, 30 and 100 μg/L) for 48 hours resulted in a dose-dependent increase of hepatic VTG expression compared to vehicle control in both
males and females. In efforts to block GPER-mediated induction of VTG expression, we co-exposed male and female SD rats with G-1 to the GPER antagonist G-6. Preliminary results were surprising as the combination of G-1/G-15 enhanced VTG expression compared to each agent singly. Overall, these data suggest that control of VTG synthesis likely involves both nuclear and membrane receptors that are sensitive to E2 activation. We are currently investigating the impacts of xenosterogens on GPER activation in control of VTG synthesis.

**1939 Study on the Subchronic Oral Toxicity of the Circuit Board Powder in Rats.**

N. Wu, Institute of Hygiene, Zhejiang Academy of Medical Sciences, Hangzhou, Zhejiang, China.

Objective To investigate the subchronic oral toxicity of the circuit board powder in SD rats. Methods Male and female SD rats were randomly divided into four groups named A, B, C and D. A and D group had 16 rats, B and C group had 14 rats, A and D group retained 4 rats for a 45d recovery experiment after the 90d subchronic experiment. A group was given normal diet as control, B, C and D groups were given mixed feed which was made from adding the circuit board powder 10, 20, 50g per kg to the normal diet. All of the groups were ate and drank freely under natural light. After 90 days' feeding, each group calculated the major organ coefficient, measured the blood biochemical parameters, and determined the content of serum triiodothyronine (T3), thyroxine (T4) and testosterone (T) after 45d, 90d and 454d recovery. Results Compared with the control group, each dose group of the circuit board powder have no significant difference in body weight. There was significant difference between the high-dose group and control group of female rats in some blood biochemical parameters. The organ coefficient of liver in the medium- and high-dose groups of female rats was also significant increase. The content of serum T3, T4 and T in each dose group were significantly higher than that of the control group after 45d and 90d, each index had the most obviously increase in the low-dose group. There was no significant difference between the high-dose group and control group in serum T3, T4 and T after 45d recovery. Conclusion subchronic oral exposure to the circuit board powder can cause liver damage in rats, and elevated the content of serum T3, T4 and T.

**1940 Activation of the Hypothalamic-Pituitary-Adrenal (HPA) Axis following Extended Exposure to Atrazine (ATR).**


While it is known that adrenal steroids impact reproduction and a variety of other physiological and behavioral functions, disruption of the HPA-axis is not typically considered in toxicological studies. Here we characterize changes in basal corticosterone (CORT) and progesterone (P4) over a 21-day exposure to the chlorotriazine herbicide ATR. Adult Wistar male rats were dosed by oral gavage with 0, 5, 25, 75, or 200 mg/kg/d given either as a single dose or multiple daily doses (7, 14, or 21 doses) at 0900 hrs ( nadir of circadian CORT rhythm) and were decapitated 30 minutes after the last dose (n=10/group). Necropsy body weight was significantly lower than controls (p < 0.05) only in the 200 mg/kg group dosed for 21 days. Adrenal weights were unaffected by ATR at any time point. Serum CORT was elevated in response to a single dose of ATR (LOEL=5mg/kg). Serum CORT was also significantly increased in the ATR groups (e.g., LOEL for one dose = 75 mg/kg, LOEL for 21 doses = 75 mg/kg). These data demonstrate a HPA response after daily doses of ATR (e.g., 75 and 200 mg/kg) for up to 21 days. Further work is needed to evaluate the effects of ATR on the circadian CORT rhythm and to discern any impact on reproductive function. This abstract does not reflect U.S. EPA policy.

**1941 2-Isopropylthioxantone (ITX) Induces Endocrine Disrupting Effects and Increases Liver Weight during Puberty in Male Wistar Rats.**

P. Hendriksen, E. Kramer, M. Groot, J. C. Rijk, R. L. Hoogenboom and A. A. Peijnenburg. Toxicology, RIKILT Institute of Food Safety, WUR, Wageningen, Netherlands. Sponsor: S. Rampaijan. 2-isopropylthioxantrace (ITX) is a photo-initiator used in the printing process of many materials. ITX has been detected as contaminant in food such as milk and fruit beverages. Previous in vitro studies revealed ITX acting agonistic on the aryl hydrocarbon receptor (AhR) and antagonistic on both the androgen receptor and estrogen receptor. In addition, ITX increases aromatase activity in vitro. To validate these findings in vivo, male rats were treated daily with 50, 150 or 500 mg/kg/day ITX during their puberty period from PND 23 to 53. For comparison, rats were treated with 50 mg/kg/day TCDD (AhR agonist), 30 mg/kg/day Casodex (AR antagonist) or 50 mg/kg/day flutamide (both AhR agonist and AR antagonist). ITX, from the lowest dose onwards, exerted a decrease in weight of the ventral prostate, and the highest dose of ITX decreased the weight of the seminal vesicles and coagulating gland, and reduced the number of spermatozoa in the cauda. The same effects, but more severe, were induced by flutamide and Casodex. Epididymal weight, caput sperm counts and preputial separation were negatively affected by Casodex and flutamide and not by ITX. ITX reduced testosterone and increased estradiol levels confirming the in vitro finding on increased aromatase activity.

500 mg/kg/day ITX-500 reduced the body weight, while it increased liver weight with 20% and relative liver weight with 38%. Whole genome mRNA expression profiling of the liver revealed a striking similarity of genes affected by ITX and flutamide. Furthermore, many TCDD-related genes, including many known AhR target genes, were regulated by ITX as well. Taken together, these results convincingly demonstrates ITX to be an endocrine disruptor that exerts AhR agonistic and AR antagonistic activities both in vitro and in vivo.

**1942 In Vivo Human Pharmacokinetics of Dibenzo(DF,EP)Chrysene (DBC) following Microdosing—Bridge the Gap between High-Dose Animal Data and Environmentally Relevant Human Exposures.**

E. P. Maden1,2, R. Corley4, K. W. Tuinelaar3, T. Ogeniben2, M. Mallat2, M. Garrard3, K. Sudalak4, T. McQuistan2,3 and D. W. Williams1,4. 1Oregon State University, Corvallis, OR; 2University of California, Berkeley, CA; 3Pacific Northwest National Lab, Richland, WA; 4Linus Pauling Institute, Corvallis, OR.

Polycyclic Aromatic Hydrocarbons (PAHs) comprise environmental mixtures of toxics resulting from the combustion of carbonaceous material, of which some high molecular weight constituents are carcinogens in animal models. There is little high molecular weight contaminants are carcinogens in animal models. There is little high molecular weight carcinogenic PAHs, especially the higher molecular weight, carcinogenic PAHs, in humans. Utilizing the sensitivity of accelerometer mass spectroscopy (AMS), 14C-labeled PAHs can be studied at environmentally relevant concentrations. Methods Twelve male and female rats, each divided into two dose groups (medium- and high-dose groups of female rats was also significant increase. The content of serum T3, T4 and T in each dose group were significantly higher than that of the control group after 45d and 90d, each index had the most obviously increase in the low-dose group. There was no significant difference between the high-dose group and control group in serum T3, T4 and T after 45d recovery. Conclusion subchronic oral exposure to the circuit board powder can cause liver damage in rats, and elevated the content of serum T3, T4 and T. 2-Isopropylthioxantone (ITX) is a photo-initiator used in the printing process of many materials. ITX has been detected as contaminant in food such as milk and fruit beverages. Previous in vitro studies revealed ITX acting agonistic on the aryl hydrocarbon receptor (AhR) and antagonistic on both the androgen receptor and estrogen receptor. In addition, ITX increases aromatase activity in vitro. To validate these findings in vivo, male rats were treated daily with 50, 150 or 500 mg/kg/day ITX during their puberty period from PND 23 to 53. For comparison, rats were treated with 50 mg/kg/day TCDD (AhR agonist), 30 mg/kg/day Casodex (AR antagonist) or 50 mg/kg/day flutamide (both AhR agonist and AR antagonist). ITX, from the lowest dose onwards, exerted a decrease in weight of the ventral prostate, and the highest dose of ITX decreased the weight of the seminal vesicles and coagulating gland, and reduced the number of spermatozoa in the cauda. The same effects, but more severe, were induced by flutamide and Casodex. Epididymal weight, caput sperm counts and preputial separation were negatively affected by Casodex and flutamide and not by ITX. ITX reduced testosterone and increased estradiol levels confirming the in vitro finding on increased aromatase activity.

500 mg/kg/day ITX-500 reduced the body weight, while it increased liver weight with 20% and relative liver weight with 38%. Whole genome mRNA expression profiling of the liver revealed a striking similarity of genes affected by ITX and flutamide. Furthermore, many TCDD-related genes, including many known AhR target genes, were regulated by ITX as well. Taken together, these results convincingly demonstrates ITX to be an endocrine disruptor that exerts AhR agonistic and AR antagonistic activities both in vitro and in vivo.

**1943 Evaluation of Tumor Pathology Concordance between Epidemiological and Rodent Studies.**

J. W. Card1, L. W. Card1, F. Kikee1, L. Haighton1, V. Lee-Brotherton1 and B. Sangster2. 1Intertek Cantox, Mississauga, ON, Canada; 2ASAT Foundation, Leidschendam, Netherlands.

To assist with evaluating human cancer risk from chemical exposures, an investigation was conducted to assess concordance at the tissue level between tumors reported in epidemiological investigations and rodent biosays for a selected group of chemicals. This investigation focused on chemicals that were considered to be probable human carcinogens (Group 2A) by IARC or “Reasonably Anticipated” carcinogens by the US NTP and for which there was evidence of tumor concordance at the organ level. The 7 chemicals identified for this evaluation, and the organ-level tumor concordances for each, were as follows: 1-amino-2,4-dibromoaniline (lung), tetrachloroethylene (liver, kidney), 1,1,-dichloroethylene (liver, kidney), inorganic lead compounds (lung, kidney, brain), 2,4,6-trichlorophenol (lymphoma), polychlorinated biphenyls (liver, gastrointestinal) and benzy1 chloride (lung). Based on pathology, concordance between rodent and human tumors for at least 1 tumor type was identified for 4 of the 7 compounds (tetrachloroethylen, trichloroethylene, inorganic lead compounds, and polychlorinated biphenyls);
however, the strength of the concordance was considered weak in all cases. For the same 4 chemicals, there were a large number of tumors which lacked concor-
dance between animal and epidemiological studies. Of the remaining compounds, insufficient data were available for an evaluation of tumor pathology (1-amino-2-4-
dibromoantraquinone and 2,4,6-trichlorophenol) or there was an absence of con-
cordance at the tissue level (benzyl chloride). In conclusion, this investigation of con-
cordance between animal and human tumors reported to be caused by specific
compounds shows that, even when chemically-induced tumors are identified to be occurring in the same organ in both rodents and humans, there is no strong evi-
dence of concordance of these tumors at the level of the affected tissue.

1944 Benchmark Dose Models for Benzene Genotoxicity Using the Diversity Outbred Mouse.
K. L. Wint1, G. E. Kisling1, D. L. Morgan1, K. R. Shockley1, D. M. Garti2, G. A. Churchill2 and J. E. French3, 1DNTP, NIEHS, Research Triangle Park, NC; 2Center for Genome Dynamics, The Jackson Laboratory, Bar Harbor, ME.
The genetic basis for inter-individual variation in response to toxicant exposure is poorly understood. The Diversity Outbred (DO) mouse is a new model that can be used to explore population based response variation. DO mice were derived from a set of eight inbred founders and are maintained by random breeding as an het-
erozygous population containing over 38 million SNPs and 5 million indels (CNV). The genetic diversity and a fine recombination haplotype structure allow high mapping resolution. In experiments reported here, J:DO male mice were ex-
posed to benzene (0, 1, 10, 100 ppm; 75 mice/group) by inhalation, 6 hr/day, 5
days/wk for 28 days in two independent cohorts (300 mice each). Samples of pre-
and post-exposure peripheral blood (PB) or post-exposure for bone marrow (BM) were evaluated for frequency of micronucleated reticulocytes (MN-RET) using flow cytometry. MN-RET showed significant increasing trends in response to ben-
zen exposure (p<0.0001) in both cohorts and were significantly higher than con-
trols in both the 10 ppm (p=0.013) and the 100 ppm (p < 0.0001) groups. Genotoxicity was highly variable within the 100 ppm groups. Linkage analysis identified quantitative trait loci (QTL) on chromosome 10 (Chr10) between megabase 26 and 35 (LOD=14.6). We identified benzene resistant and susceptible Chr10 QTL genotypes from the lowest and highest quintiles of the BM MN-RET response in the 100 ppm group. Animals in the 0, 1, and 10 ppm groups having these genotypes were placed into the resistant or susceptible groups for modeling. The benchmark concentration (BMC) and its 95% lower confidence limit (BMCL) were estimated from the best-fit model for the most resistant (BMCL > 5.9 ppm) and most susceptible (BMCL < 0.01 ppm) animals, a 590-fold difference between subpopulations. These data suggest that an uncertainty factor of 100 for interspecies and intraspecies variation may not be sufficient for calculation of a human benzene reference concentration based on benzene genotoxicity.

1945 Is There a Subset of Susceptible Individuals in Controlled Air Pollution Studies?
M. Seeley and J. E. Goodman, Gradient, Cambridge, MA.
To establish primary National Ambient Air Quality Standards for criteria air pollu-
tants such as nitrogen dioxide (NO2), ozone (O3) and sulfur dioxide, US EPA eval-
uates results from epidemiologic controlled human exposure, and animal studies. For the controlled human exposure studies, which focus on individuals with asthma, there is concern that there may be a subset of susceptible individuals who respond to lower concentrations than the average individual with asthma, but whose response is obscured by evaluating group-level data. To address this, we iden-
tified controlled human exposure studies with more than one exposure concentra-
tion that provided individual-level data for airway hyper-responsiveness. We identi-
fied three studies involving exposure to NO2 (including data for 38 subjects), and one study each for ozone (O3) and sulfuric acid (each including data for 10 sub-
jects). For each study, we compared individual data across exposures to determine whether there were individuals who were consistently more or less responsive, de-
fined by having a response that differed by more than two standard deviations from
the group mean response. To determine whether evaluating group-level responses
might obscure responses for potentially susceptible individuals, we compared the concentration-response function (CRF) for all subjects to the CRF for a subset that excluded less responsive subjects and the CRF for only more responsive subjects. Among the five studies, we identified only one subject, in one study, who was con-
siderably more responsive at both exposure levels in the study. The shapes of the
CRFs, however, were similar for the group as a whole and the more responsive sub-
ject. We also identified one subject, in one study, who was less responsive at all ex-
posure levels in the study. Exclusion of data for this subject did not qualitatively change the shape of the CRF. Our analysis of these five studies indicates that evalu-
ating group-level responses to air pollutants does not obscure CRFs for potentially
susceptible individuals.

1946 Concordance of Transcriptional and Apical Benchmark Dose Levels for Conazole-Induced Liver Effects in Mice.
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The ability to anchor chemical class-based gene expression changes to phenotypic
lesions and to describe these changes as a function of dose and time can inform
mode of action and improve quantitative risk assessment. Previous research identi-
fied a 350-gene cluster commonly responsive to three hepatotumorigenic conazoles (cypro-, epoxi- and propiconazole). Among these conazoles (tri-
adinefon and myclobutanil), the present assessment encompasses four tumorigenic and one non-tumorigenic conazole. Transcriptional benchmark dose levels (BMDL) were estimated for a subset (~50 genes) of the gene cluster that had a ≥5
fold change in signal intensity at the tumorigenic dose and demonstrated dose-re-
sponsive behavior. The modeled genes primarily encompassed pathways involved in
Phase II metabolism and biliary excretion (such as Gst3, Gst3, Abcc3, Akr1b7), mirrored each conazole's tumorigenic potency. The 30-day BMDL for liver tumors. Discordant 30-day BMDL and BMDL estimates were obtained for the non-
tumorigenic conazole (myclobutanil). Potency differences seen in the dose-re-
sponsive transcriptional mapping of some of these biomarker genes, particularly those involved in Phase II metabolism and biliary excretion (such as Cyp2b10, Lcn13, Abcc4, Akr1b7), mirrored each conazole's tumorigenic potency. The 30-day BMDL and BMDL estimates corresponded to tumorigenic potency on a mg/kg-day basis with cyproconazole > epoxiconazole > propiconazole > triadimefon > myclobutanil (non-tumorigenic). These initial results support the utility of measuring short-term gene expression changes to inform quantitative risk assessments from long-term exposures. This abstract does not reflect EPA policy.

1947 Assessment Factors for Susceptible Populations—Analysis of Airway Response during Short-Term Exposure to Volatile Chemicals.
M. Johansson, G. Johanson and M. Öberg, Institute of Environmental Medicine, Stockholm, Sweden.
Health-based guidance values for short-term exposure to airborne hazardous chem-
icals are developed to support the protection of the general population, including sus-
cetable sub-groups such as asthmatics, in the case of sudden release of chemi-
cals. The Acute Exposure Guideline Level (AEGL) program is one of the most well-
known set of short-term values. Our analysis of AEGL documents reveals that only
8% include data on asthmatics. A comparison of documents in nine additional sets
of short-term values shows that data on asthmatics are frequently disregarded. The
aim of the present study was to investigate the experimental support of a general
difference in airway response between healthy and asthmatic individuals during
short-term exposure. We performed a review of experimental data from 108 studies
including both asthmatic and healthy subjects exposed to airborne chemicals dur-
ing identical experimental conditions. In total, experimental data for 19 chemicals
and 9 mixtures were identified. Thresholds for airway response and the difference
between asthmatic and healthy subjects were calculated in each study. The differ-
ence between subgroups was compared to the general assessment factors applied
for susceptible populations in the derivation of guidance values. In addition, dose-re-
sponse relationships of four high volume chemicals were calculated for healthy and
asthmatic individuals, separately, to identify threshold concentrations for effects on
lung function. Our results show that asthmatic individuals generally are more sus-
cetable than healthy individuals. An inter-individual assessment factor of three
may not be sufficiently protective for all chemicals.

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Chemicals and heat exposure are common hazards found in the workplace and
their coexposure may lead to increased health risk due to potential interactions:
some chemicals can affect the thermoregulatory mechanisms and reduce the
worker's capacity to adapt to heat, while heat exposure can modify physiological parameters in various parts of the body. Therefore, the aim of this research was to identify the occupations at risk of concomitant exposure to heat and chemicals in Quebec (Canada). To achieve this goal, a list of occupations from all industry sectors of Quebec was built from national employment data, and occupations at risk of heat stress were determined with a risk rating matrix based on the probability of occurrence and the severity of health effect. For occupations judged as having significant and critical heat stress risk, the presence of chemicals that may be found at work was documented (solvents, dust, pesticides, polycyclic aromatic hydrocarbons, toxic gases, heavy metals, asbestos/silica and reagents/other chemicals) and a list of occupations with high potential of simultaneous exposure to heat and chemicals was produced and submitted to a panel of experts for prioritization. From an initial set of 1010 occupations, 257 were judged as having significant and critical heat stress risk and 136 of these were included in the list of occupations with high potentially simultaneous exposure. Experts identified 22 priority occupations grouped in metal manufacturing, non-metallic mineral product, construction and public administration sectors. This innovative approach, based on a matrix analysis and judgment of experts, served as a useful tool to identify sectors/occupations where health risk assessments, preventive interventions and regulatory practices should be conducted or developed.

**1949 Acute and Chronic Noncancer Inhalation Toxicity Factors for Acrylonitrile.**

J. Lee, R. L. Granzi and S. Shirley. Toxicology, TCEQ, Austin, TX.

Acrylonitrile (AN) is used extensively in the production of plastics, synthetic rubber, nitrite elastomers, resins, and acrylic fibers. The USEPA indicates that Texas contributes 11% of the nations reported ambient AN emissions annually. Inhalation of AN vapors can cause respiratory irritation, and at higher levels, neurological symptoms including dizziness, weakness, headache, and impaired judgment. To ensure that the general public in Texas is protected against potential inhalation effects from AN exposure, the Texas Commission on Environmental Quality (TCEQ) has developed acute and chronic reference values (ReVs). An acute ReV (1-hr exposure duration) of 1100 μg/m3 was derived based on no signs or symptoms observed in human volunteers exposed to AN for up to 8 hours. A chronic ReV of 2.2 μg/m3 was derived based on benchmark dose modeling for increased nasal lesions observed in female rats. The chronic ReV is comparable to the California EPA reference exposure level of 5 μg/m3. Effects Screening Levels (ESLs) were calculated from ReVs by applying a target hazard quotient of 0.3, to account for possible cumulative exposure. ESLs are used to evaluate modeled ground level concentrations due to emissions from facilities during air permit reviews. The corresponding acute and chronic ESLs were 350 and 0.7 μg/m3, respectively. Reproductive/developmental animal and epidemiological data were not used to derive ReVs since AN is not expected to be a developmental or reproductive toxicant in the absence of significant maternal toxicity. Furthermore, the overall carcinogenic weight-of-evidence shows that while AN is capable of causing tumors in rats and mice at high doses, AN does not appear to contribute to the development of cancerous tumors in humans. Thus, no inhalation unit risk factor was derived. The derived chronic ESL, however, is within the range of the concentrations at 1 x 10-5 cancer risk estimated by USEPA and thus, is expected to be protective against potential cancer risk.

**1950 Hexavalent Chromium Carcinogenicity: Use of a Nonlinear-Threshold Assessment to Develop a Cancer-Based Chronic Inhalation Reference Value.**

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It is important to conduct up-to-date chemical assessments for known human lung carcinogens such as hexavalent chromium (CrVI). An updated carcinogenic assessment has been conducted for CrVI, which has been the subject of recent scientific debate. In addition to default linear low-dose extrapolation methods used to calculate an inhalation unit risk factor (URF), the study authors believe epidemiological data supported by data relevant to the mode of action (MOA) are sufficient to justify considering the results of a nonlinear-threshold carcinogenic assessment for comparison to URF-based de minimis excess risk (e.g., 1 in 100,000) air concentrations. The intent of the current study is not to perform an exhaustive weight of evidence evaluation of all data potentially relevant to the MOA (or MOAs), but rather to present available summary MOA information and statistical evidence interpreted as supporting a potential practical threshold for CrVI-induced carcinogenicity and the results of the consequent nonlinear-threshold inhalation carcinogenic assessment. Relevant epidemiological studies available in the scientific literature were reviewed and additional statistical dose-response analyses conducted to interpret the practical thresholds and points of departure (PODs). Occupational-to-environmental dosimetric adjustment of the “sub-threshold” cumulative exposure POD selected (0.195 mg CrVI/m3-yr) resulted in a POD16 of 0.0071 mg CrVI/m3. Uncertainty factors (total UF of 30) were then applied to derive a cancer-based chronic inhalation reference value (ReV) of 0.24 μg CrVI/m3.

**1951 Hypothesis-Based Weight-of-Evidence Evaluation of the Human Carcinogenicity of Toluene Disosyanoate.**

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Humans are exposed to toluene disosyanoate (TDI) primarily through inhalation in workplaces where it is used or produced and is classified as a possible human carcinogen, based primarily on increased tumor incidences in rodents treated with TDI by oral gavage. We used the hypothesis-based weight-of-evidence (HBWoE) method to conduct a novel evaluation of all the evidence regarding the hypothesis that TDI is a human carcinogen. We weighed the available data from epidemiology, animal toxicology, and mechanistic studies in terms of quality and relevance to humans, allowing each data set to inform one another. We then evaluated all of the data together to determine whether TDI carcinogenicity is plausible in humans. Our analysis determined that the epidemiology data are not sufficiently robust to support TDI as a human carcinogen; the few positive associations are more likely attributable to alternative explanations than causation. The experimental animal studies indicate that inhalation exposure to TDI does not induce tumors in rats or mice. Tumors observed after oral gavage exposure are most likely due to the conversion of a small amount of the administered TDI to toluene diamine (TDA), which also causes tumors in rats and mice when administered through this physiologically route. Further, TDA is formed in vivo TDI genotoxicity assays. While TDI is genotoxic in these assays, it is not genotoxic in rodents or humans in vivo after inhalation exposure (during which TDA is not formed). Our HBWoE analysis indicates that the conversion of TDI to TDA is necessary for carcinogenicity to occur, but it does not occur in mammalian species under physiological exposure conditions. Thus, our analysis demonstrates that a causal association between TDI exposure and carcinogenic effects is not plausible in humans. This analysis not only provides insight about TDI carcinogenicity, but it addresses the larger issue of combining human, animal, and mechanistic data for risk assessment.

**1952 Evaluation of Toxic Effects of 3-MCPD and Associated Esters in Rats.**

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3-Monochloropropene-1,2-diol (3-MCPD) is regarded as a rodent renal and Leydig cell carcinogen. Furthermore, 3-MCPD esters may be generated in various foods and food ingredients as a result of food processing. Since there are limited reports about toxicity of these compounds, we conducted the following two studies: 1) In vivo genotoxicity study (B6AF1 and B6AF1 gpt gene mutation assays in the kidneys and testes, micronucleus assay) with male F344 gpt delta transgenic rats carrying reporter genes for mutations, treated by gavage with a carcinogenic dose of 3-MCPD (3.6×10^-4 mol/kg body weight) or its esters, i.e., 3-MCPD palmitate diester (PD), 3-MCPD palmitate monoester (PM) and 3-MCPD oleate diester (OD) at the molar equivalents for 4 weeks. 2) 13-Week subchronic toxicity study of these compounds with F344 rats at three doses where the highest was the same as that in the genotoxicity study. In the first study, no clear in vivo genotoxicity of 3-MCPD or its esters was observed in the three assays. In the 13-week study, the absolute and relative kidney weights were significantly increased in rats treated with 3-MCPD at a carcinogenic dose and the esters at high and medium doses. Relative liver weights were significantly increased in the 3-MCPD-treated rats and the high dose ester-treated rats, except for female rats treated with PM. By 4 weeks, 1 male and 5 female males of 3MCPD group died from the renal tubular necrosis. On histopathological analysis, a significantly increased incidence of renal mineralization in all high dose ester-treated females and of apoptotic epithelial cells in the epididymis of 3-MCPD and high dose ester-treated male were observed as compared to vehicle control. The results suggest 3-MCPD and its esters may be non-genotoxic but have potential toxicity for kidneys and epididymides of rats. NOAELs of PD, PM and OD were suggested to be 14, 8 and 15 mg/kg/day, respectively.
1953 A Simulation Study Investigating the Risk of Obtaining BMDs Which Are Higher Than the "True" BMD.


A central aspect of quantitative health risk assessment of chemicals is the derivation of the point of departure (POD). Historically, calculating NOAELs has been the most common way to obtain the POD. However, Benchmark Dose (BMD) analysis has been recommended as a more robust alternative, as it uses data from all dose groups in the study to derive the POD. An important step in the BMD approach is the identification of the BMDL, i.e., the lower limit of the confidence interval for the BMD. The aim of this study was to investigate how often the estimated BMDL corresponds to an effect greater than the critical effect, thereby underestimating the risk in the derivation of guidance values.

In this simulation study we assume a known sigmoidal dose-effect curve (and hence a known BMD) and examine how often the BMDL is higher than the "true" BMD. Effects were generated by Monte Carlo simulations using 5 different dose placement scenarios. All scenarios used 4 dose groups (control, low, medium and high dose) and logarithmic dose spacing. Each scenario was simulated using 5, 10, 20 or 50 animals per group, assuming dose-effect curves typically seen in continuous data from experimental studies. The BMD and the BMDL values were calculated using a set of 5 nested exponential models implemented in available BMD-softwares (BMDs and PROAST).

The results suggest that the risk of obtaining a BMDL which is higher than the true BMD can be considerably larger than normally expected. In one scenario the BMDL was higher than the true BMD in 74% of the simulations. This occurs because the second model in the nested set of models lacks the ability to level off at higher doses. It is therefore suggested that the second model in the nested set (γ=a*e^b) should be used with caution. It is also important to visually inspect the dose effect curve and the individual data points.

1954 Characterizing the Impacts of Uncertainty and Scientific Judgment in Exposure Limit Development.


There is a misperception by some that exposure limits are precise estimates. In the eyes of risk managers, one discrete value is often considered to be "correct" and all others considered "incorrect." Exposure limits should be evaluated based on whether the value is derived in a manner "consistent with current principles" or "not consistent." An analysis of current risk assessment methods was conducted to identify the bases for variability in exposure limits for individual chemicals. The role of scientific judgments, risk policy perspectives, and evolving scientific methods were evaluated in the context of exposure limit setting methods. A systematic methods analysis shows that important drivers to be considered in evaluating acceptability of an exposure limit include: thoroughness of the review of available data, interpretation of results according to current scientific principles under the regulatory framework being used, and consideration of sufficient sources of variability and uncertainty. Sources of variability that may be encountered in risk assessments performed by different industrial hygienists or toxicology professionals using identical data sets include: selection of the point of departure, uncertainty factors used for data extrapolation, and use of adjustments for toxicokinetics, among others. These and related considerations form the basis of a "quality evaluation" process proposed for assessing the robustness of an exposure limit. Transparency in methods to assure robustness is a core principle embedded in risk assessment methods harmonization. Application of a systematic quality evaluation process provides for more informed use of exposure limits for risk management. A clear understanding of the basis for disparate values can provide useful information regarding the current level of uncertainty in the science and the level of confidence appropriate in using different exposure limits to characterize risk.

1955 Chemical-Induced Methemoglobin Formation and Exploration on Its Biological Threshold.

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Methemoglobin (metHb) is hemoglobin with the iron oxidized from Fe2+ to the Fe3+ state. An accumulation of metHb in the blood (methemoglobinemia) is often observed in response to exposure to many chemical agents, such as aniline and aniline derivatives like dapsone, and is often used as the basis for the derivation of non-cancer risk values. Therefore, an understanding of the relative sensitivity of common laboratory animals compared to humans to these metHb-forming agents is important. In this research, we compared the relative sensitivities to dapsone, a metHb-forming agent in rats, mice, and humans based on the data from in vitro and in vivo studies. In vivo data indicate that humans are more sensitive to dapsone than rats, followed by mice. However, in vitro results are inconsistent in terms of metHb formation. The inconsistency can be explained by differences in liver metabolism among species. In vitro comparisons between the parent compound (dapsone) and its metabolite (hydroxylamine) also suggest that the metabolite is more potent than the parent compound to induce metHb. Due to a higher sensitivity in vivo, the rat might be a more suitable animal model than the mouse for predicting metHb-forming effects in humans. In addition to the relative sensitivities, we further examined the background levels of metHb to explore the potential for identifying the biologically significant threshold of metHb formation after exposure to metHb-forming agents. F344 rats were used as an example. We identified a 100% increase above the control mean as a benchmark response (BMR) based on the collected metHb background levels. A comparison of the identified BMR and one previously derived from the control mean, as commonly used in benchmark dose modeling, suggests that the new BMR generates lower or comparable benchmark dose lower confidence limits (BMDLs). Therefore, a 100% increase above the control mean could be another way to establish a BMR for methemoglobinemia in F344 rats. The views expressed in this abstract are those of the authors and do not necessarily reflect the views or policies of the U.S. EPA.

1956 Analysis of Possible Changes to the Levels of Concern for Polycyclic Aromatic Hydrocarbons in Seafood.


Anthropogenic contamination of coastal regions with oils from drilling operations, spills and tanker leaks has impacted coastal communities and seafood safety for decades, and will likely continue based on our oil-based economy. Federal and state agencies suspend commercial and recreational harvests in oil-affected regions based on testing for petrogenic toxics, including polycyclic aromatic hydrocarbons (PAHs). Criteria for re-opening affected fisheries are dependent on human health risk-based levels of concern (LOCs) developed by the US Food and Drug Administration (FDA) for specific petroleum-related contaminants in seafood, including seven carcinogenic PAHs. As with the Deepwater Horizon oil spill and other oil spill events, FDA cancer-based risk assessment for seafood consumption applies the US Environmental Protection Agency (EPA) cancer slope factor for benzo[a]pyrene (BaP), and relative potency factors (RPFs) for the remaining six PAHs relative to BaP. Use of LOCs in risk assessment is based on numerous assumptions. Here we examine how LOCs for PAHs change with modifications based on: (1) variable seafood consumption rates using data reflective of high-end seafood consumers in affected regions, (2) an expanded list of carcinogenic PAHs and changes in relative potency factors as described in the EPA draft document “Development of a Relative Potency Factor Approach for Polycyclic Aromatic Hydrocarbon Mixtures,” (3) the proposed change in the EPA cancer slope factor for BaP (the index carcinogenic PAH), and (4) exposure assumptions relevant to children, including incorporation of an age-dependent adjustment factor for PAH carcinogenesis. Analyses provide an indication of both direction and magnitude of changes in LOCs associated with each possible modification. This study suggests that LOCs for PAHs in seafood from the Gulf of Mexico may be more numerous and markedly lower than the suite currently applied relative to the Deepwater Horizon oil spill.

1957 The Use of Genetically Modified Mice in Cancer Risk Assessment: Challenges and Limitations.


The use of genetically modified (GM) mice to assess carcinogenicity is playing an increasingly important role in the safety evaluation of chemicals. While progress has been made in developing and evaluating models such as the Tpr53+/−, the Tg.AC and the raf32 models, the suitability of these models as replacements for the conventional rodent cancer bioassay and for assessing human health risks remains uncertain. The objective of this research was to evaluate the prospective use of GM mice and the recently developed accelerated cancer bioassays in evaluating the potential health risks associated with exposure to carcinogenic agents. We compared the published results from the GM bioassays with those obtained using the National Toxicology Program's conventional chronic mouse bioassay for their potential use in risk assessment. To date, the GM models have shown moderate success in distinguishing carcinogens from non-carcinogens. Analysis of information
from different studies indicates that the GM models are less efficient in detecting carcinogenic agents but more consistent in identifying non-carcinogenic agents. We identified several issues of concern related to the assay design of GM models (e.g., sample size, study duration, genetic stability, and reproducibility) as well as pathway-dependent effects, and different carcinogenic mechanisms operable in GM and non-GM animals. The use of the GM models for dose-response assessment is particularly problematic as these models are, at times, much more or less sensitive than the conventional cancer bioassays. Thus, the existing GM mouse models may be useful for hazard identification, but will be of limited use for dose-response assessment. Hence, caution should be exercised when using GM mouse models to assess the carcinogenic risks of chemicals. 

Disclaimer: The views expressed in this abstract are those of the authors and do not necessarily reflect the views or policies of the USEPA or the NIEHS.

1958 Integrating Local Communities in the Health Risk Assessment Process following the Deepwater Horizon Oil Spill—A Focus on Vietnamese-Americans.


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Vietnamese-American populations in southeast Louisiana consist largely of commercial and subsistence fisherfolk and represent one of the highest seafood consuming groups in the Gulf South. We collected targeted survey data from a Vietnamese-American fishing community in Orleans Parish Louisiana, to determine shrimp consumption rates, body weights, ages, and genders in order to conduct deterministic and probabilistic health risk assessments tailored to this unique population. Cancer risk estimates and levels of concern (LOCs) for several poly- cyclic aromatic hydrocarbons were determined using GC/MS in SIM mode on a sample of white shrimp collected from the Gulf with shrimpers from this community. Our approach also developed health risk assessments using LOCs, oral slope factors, and risk levels used by the Food and Drug Administration and the Natural Resources Defense Council. Our results demonstrate the need to include key populations in the risk assessment process and measure risk model parameters in such populations rather than rely solely on generic exposure assumptions.


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There is a growing trend of using botanical raw materials in personal care products, such as those used in traditional medicine or those from the many exotic plants which may be part of local folklore or food use. These materials present a challenge to both the Product Developer and Regulator alike, in ensuring the development of safe cosmetic products, from the complex composition of the materials through to the lack of documented data on history of use and safety. This paper presents a strategy for the safety assessment of these botanicals used in cosmetics, based on various risk assessment concepts. A critical first step in the risk assessment is the characterisation of the botanical raw material, with key measurements being identified. A complete understanding of “what is known” about the material should then be developed from both literature sources as well as traditional knowledge. It may then be possible to determine whether a history of safe use (HOSU) can be established and if the material can be considered safe at that stage for particular cosmetic applications. If not then further risk assessment approaches (e.g comparative approach, threshold of toxicological concern (TTC)) are proposed. Finally, in order to complete the risk assessment there may be a need to fill gaps in the hazard profile and/or potential consumer exposure scenarios, by conducting some further testing.

1960 Simulation of Acute Reference Dose (ARD) Setting for Pesticides in Japan.


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We conducted simulations of Acute Reference Dose (ARD) setting, based on review documents of 208 pesticides published by Food Safety Commission (FSC) in Japan. These pesticides were evaluated in FSC during last 8 years. We applied the conceptual framework of Solecki, et al.(2005) to create and implement a conceptual framework adapted to current assessment needs in Japan. Through this process, we were able to set the ARDs for over 90% of those 208 pesticides. The studies that provided the rationale for ARD setting were primarily reproductive and developmental toxicity studies, acute neurotoxicity studies, and pharmacology studies. It was not necessary to establish ARDs for approximately 30% of the pesticides simulated in the present study. Some of the simulated ARDs might be conservative, and some endpoints for ARD setting might not be proper, because the published data obtained in the present study were written for acceptable daily intake (ADI) setting. We were unable to set an ARD for 14 pesticides because of insufficient data on acute toxicities. This could be improved by more complete record-keeping (for example, the type of changes observed immediately after administration, and the duration of the observation period). Furthermore, we categorized the 208 pesticides by mechanism of action or chemical structure. In comparison of absolute ARDs or relative ARDs to ADI among the categories, considerable number of pesticides with similar mechanisms of action or similar chemical structure also showed similar ARDs.

1961 Tumour and In-Life Data from CD-1 Mouse Dietary and Oral Gavage Tumorigenicity Studies, Completed over the Period of 1995 to 2011.

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A previous review (1) of Crl:CD-1 ® (ICR) BR mouse tumorigenicity studies indicated terminal (Week 104) mortality values of 5±10.2% for males and 7.7±7.7% for females, with no major differences between the dietary and oral gavage routes of administration. Expanding on this, the aim of this review was to establish if the in-life data and tumour profile had changed over time and if there were any differences between the routes of administration. Data was analysed from over 60 mouse studies terminated at or about 2 years and completed over the period of 1995 to 2011. Analysis of bodyweight gain and food consumption over the first year revealed lower values for oral gavage studies (both sexes) when compared with dietary studies. The analyses over time (Period 1, 1995-2001 against Period 2, 2002-2011), performed for oral gavage studies only, did not show any remarkable differences. The most prevalent tumours were hepatocellular adenoma/carcinoma and adrenal adenoma (males), bronchial/alveolar adenoma/carcinoma, Harderian gland adenoma, and haemopoietic tumours (males and females), and uterine polyps (females). Generally, excluding the latter, these tumours were also the major pathological factors contributory to death (FCTD). Other major FCTD included urinary tract lesions (males) and ovariian haemorrhagic cysts (females). The most prominent difference between dietary and oral gavage studies was a lower incidence of hepatocellular tumours (males) in the latter. Analyses over time (for oral gavage studies only), indicated a lower incidence of hepatocellular tumours (males) and Harderian gland adenoma (both sexes) in Period 2. In conclusion, although there were some differences in the mortality patterns, in-life parameters, tumour profile and FCTD between the alternative oral routes and between the time periods, the CD-1 mouse has a well-defined tumour profile and continues to be suitable strain for use in tumorigenicity bioassays.

1962 Factors Influencing the Outcome of Developmental Neurotoxicity Studies of Bisphenol A—Implications for Regulatory Testing and Risk Assessment.

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Developmental neurotoxicity (DNT) of bisphenol A (BPA) has been investigated in a large number of studies. However, there are discrepancies in the results reported between these studies. This investigation aims to identify and analyze factors that have contributed to these differences and to assess whether there are sex-differences in the sensitivity of certain endpoints or tests used in DNT-studies.

Forty-four DNT studies of BPA were identified in the open literature. Details about study design and results from each study, as well as the criteria for DNT testing according to the standardized OECD test guideline (TG) 426, were collected in a data base. This enabled systematic and detailed comparisons between studies as well as to the TG criteria. Multivariate analyses of the compiled data were applied in order to investigate how different factors of the study design contribute to differences in outcomes between studies.

The results from this investigation indicate that the choice of behavioral endpoints and the behavioral test paradigms used seem to have the largest impact on study outcome. In general, the most sensitive endpoints were the ones not required according to the standard OECD TG 426. Interestingly, non-standard endpoints seem to be especially important to detect effects in females.

One main conclusion from this investigation is that non-standardized studies seem to provide information that could be pivotal for the risk assessment of BPA, especially in the identification of effects in females. There is a need to develop tools that improve the usability of non-standardized studies in risk assessment of chemicals, particularly with regard to endocrine disrupting compounds, and that can facilitate the evaluation of data obtained from such studies.

1963 Development of Human Health Benchmarks for Pesticides.

S. Ramasamy1, E. Doyle1, B. Behl1, B. May2, M. Goodis2 and S. Knizner3.

On March 22, 2010, EPA Administrator, Lisa P. Jackson announced a new drinking water strategy that outlines four principles to expand public health protection. One of these principles is to use the authority of multiple laws to more effectively protect drinking water, by sharing data collected under different statutes. Under this principle, the Office of Water (OW) and the Office of Pesticide Programs (OPP) collaborated to develop the human health benchmarks for approximately 350 pesticides (HHBPs) that are currently registered to be used on food crops. These HHBPs are levels of certain pesticides in water at or below which adverse health effects are not anticipated from one-day or lifetime exposures. EPA derived the HHBPs using toxicity studies conducted on laboratory animals submitted to EPA under the Federal Insecticide, Fungicide and Rodenticide Act (FIFRA), the Federal Food Drug and Cosmetics Act (FFDCA) as amended by the Food Quality Protection Act (FQPA), with the typical methods used for developing drinking water health advisories under the Safe Drinking Water Act (SDWA). Specifically, EPA used the acute or chronic reference doses established for human dietary assessment for the most sensitive population and applied standard exposure assumptions used in calculating drinking water health advisories based on gender, age, body weight, and water consumption of target populations. For the acute benchmark, the entire exposure is assumed to occur from drinking water (100 percent). For the chronic benchmark, EPA applied a default relative drinking water source contribution (20 percent), assuming additional exposure may arise from other sources. The HHBPs are not enforceable. EPA is providing the benchmarks for informational purposes for use by states, water systems and the public to help understand monitoring data for pesticides that have no drinking water standards or health advisories.

The HHBPs can be found on EPA’s website at: http://www.epa.gov/pesticides/hhbp

1964 Differences between US EPA Iris Inhalation Reference Concentrations (RfCs) and European Chemicals Agency (ECHA) Long-Term Inhalation Derived No Effect Levels (DNELs) for the General Population.

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ECHA previously released manufacturer/importer REACH (Registration, Evaluation, Authorization and Restriction of Chemicals) chemical registration files, making available the Derived No Effect Levels (DNELs) for hundreds of substances. The DNEL is defined as “the level of exposure above which humans should not be exposed” and needs to reflect the likely route(s), duration and frequency of exposure. Typically, DNELs are developed by Industry using ECHA’s guidance; the registrant has the final decision on the selection of the key studies, endpoints of concern, and the assessment factors to account for sources of uncertainty, instead of a regulatory agency such as the U.S. EPA for the IRIS program. Considering this major difference, we have conducted a comparison analysis between ECHA’s long-term inhalation DNEL for non-carcinogenic effects for the general population and IRIS’ chronic RfC, which is defined as “an estimate (with uncertainty spanning perhaps an order of magnitude) of a continuous inhalation exposure to the human population (including sensitive subgroups) that is likely to be without an appreciable risk of deleterious effects during a lifetime.” Overall, despite being defined similarly, many of the current U.S. EPA IRIS inhalation RfCs for which long-term inhalation DNELs were available are lower by a factor of at least 10. The differences between these RfCs and DNELs may be a result of differences in the processes by which they are derived, including differences in the availability of toxicity data, as unpublished data or ECHA-required data may be used in the REACH assessments [ECHA requires the registrant to conduct toxicity testing (when data for a required endpoint are not available)], the entity (industry versus regulatory agency) responsible for generating the reference values, the methodologies and selection of default values used in the derivation, and the process by which the final values are accepted.

1965 Assessment Factors in Human Health Risk Assessment and Their Associated Level of Safety.

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Human health risk assessment requires solid information on adverse effects after long-term exposure. Because of ethical considerations, human data or long-term studies with animals are in general scarce. Consequently, reliable assessment factors (AF) are needed in risk assessment to overcome differences between short-term animal studies and the human situation. In human health risk assessment, traditional AF are established, e.g. a factor of 100 (10 x 10) for interspecies/intraspecies extrapolation. Some of these AF were substantiated by either deterministic or probabilistic approaches, whereas other factors lack a scientific rationale. Furthermore, the level of safety achieved with the application of more than one AF has remained an open question so far. Our analyses aimed therefore at the derivation of AF and their corresponding level of safety when applied solely or in combination in human health risk assessment. The toxicological database RepDose(TM) (containing appr. 2500 repeated dose studies on appr. 700 chemicals) served as the basis for the derivation of AF. Pairs of NOEL ratios - e.g. obtained from two studies with the same chemical and species, but different application durations if time AF were analyzed - were plotted in a distribution curve. Distribution curves for interspecies and time extrapolation were evaluated. All distribution functions were best described by lognormal curves. Furthermore, the combinations of these distributions were analyzed in probabilistic approaches by means of Monte Carlo simulations. On the basis of the resulting distribution curves, overall AF were determined, for which the corresponding percentile indicates the associated level of safety. Our results demonstrate that the probabilistic approach is well suited to derive AF, provided that the database is comprehensive and contains studies of good quality. RepDose: Trademark of the Fraunhofer Gesellschaft zur Foerderung der ange-wandten Forschung e.V., 80686 München, DE.
Multiple studies have noted that many carcinogenic polycyclic aromatic hydrocarbons (PAHs) also demonstrate immunosuppressive effects. Here, we have tested the strength of this association by deriving both cancer and immunotoxic relative potency factors (RPFs) for multiple PAHs, using the well-studied carcinogenic and immunotoxic compound benz(a)pyrene (BaP) as the index PAH. Our quantitative analysis demonstrated a correlation between the immunosuppressive and carcinogenic potential of PAHs. As evocation of immune destruction has been identified as an "emerging hallmark of cancer" (Hanahan and Weinberg, 2011 Cell 144: 646-674), we propose that this association between carcinogenic and immune suppression activities of PAHs offers an opportunity to improve cancer risk assessment for PAH mixtures. While current assessments of cancer risks from PAH-containing fractions of environmental mixtures rely on a component-based RPF approach, we hypothesize that a more scientifically defensible whole mixtures approach would be advanced not only by a battery of inexpensive, short-term tests of different modes of carcinogenic action (e.g., genotoxic and non-genotoxic), but also a battery of tests for immunosuppressive activity. The addition of immunotoxicity tests may improve the ability to discern the similarity of an untested environmental mixture with reference mixtures that have cancer slope factors. Therefore, we performed a comprehensive review of the immunotoxic effects of BaP, and developed an integrated knowledge map illustrating immunosuppressive effects of BaP and, when known, tissue-specific mechanistic information. Our analysis revealed that research to date has not yet identified a transcriptional signature(s) for immune suppression to aid in the development of a specific battery of short-term in vitro test for immunotoxicity. Thus, future directed research to develop and validate a battery of tests for immunotoxicity is warranted and could find widespread use in assessing cancer risk from environmental PAH-containing mixtures.
deriving safe exposure levels for these materials. Each of these materials comes in a multitude of forms, varying in size, shape, structure, purity, and surface coating, which affect chemical properties, such as solubility, reactivity, and propensity for agglomeration. All 3 nanomaterials have multiple repeated exposure inhalation studies available, suitable for use in dose-response assessment, but only nAg has repeated dose oral studies, and none of the materials has a suitable dermal study. The lung was the critical target in the inhalation studies of all 3 materials, with pulmonary inflammation being the most sensitive endpoint. For nAg, the liver was also identified as a target for both inhalation and oral exposure (bile duct hyperplasia). Using these data, it is possible to derive safe exposure levels for long-term exposure to the specific forms of nAg, nTiO2, and CNT tested in these studies. It is, however, unclear to what extent these levels may apply to other forms of nAg, nTiO2, and CNT. Mechanistic data for all 3 nanomaterials show that the specific physical and chemical characteristics of the particles can influence toxicity in defined ways. However, the relationships are complex, and as yet, only partially understood, making the task of estimating safe levels of a particular nanomaterial challenging. Toxicity assessments for nanomaterials must take these additional uncertainties into account. (The views expressed in this abstract are those of the authors only and do not necessarily reflect the views or policies of the U.S. CPSC).

**1971 Quality Assurance and Quality Control for an xCelligence High-Throughput Risk Assessment Assay.**

M. Stampp1, F. Ackah1, V. Charoenruk1, T. Pan2, C. Jin1, Y. A. Abassi1, X. Xu1, X. Wang1, B. Huang1, D. Kinniburgh1, W. Zhang2 and S. Gabos3.

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Taking advantage of impedance based non-invasive, real time cell analysis technology, and human cell lines with different organ origins, a High Throughput Risk Assessment (HTRA) assay is under development to meet 21st century testing requirement for hazard identification, chemical mode of action understanding, and eventually human health risk assessment. The HTRA assay is run on an fully automated system, composed of 4 xCELLIGENCE RTCA HT stations running 384 well EPlates, liquid handler, and compound/cell culture hotels. Impedance signal derived from adherent cells in the EPlate 384 are continuously monitored for 100 hours throughout the treatment and response to testing chemicals. To assess and validate the usefulness of this new in vitro screening system for identifying general toxicity of environmental hazards, 14 chemicals with known in vivo LD50 values were tested with 11 different concentrations in duplicate or quadruplicate, and repeated in 2 separate experiments. The quality assurance program includes the generation of SOPs for experiments and data analysis. Specifically, experimental plate layout to reduce false responses is discussed. The quality control system includes variability and reproducibility analysis. Coefficient of variation, signal to noise ratio and Z factor were calculated to evaluate variability. The analysis excluded one cell line from the cell panel. Intra-plate reproducibility of impedance signals was within acceptable limits. Intra-plate reproducibility of IC50 values from data collected on LC50 values from data collected on cell lines, was acceptable. Inter-laboratory validation is in progress. In summary, results from HTRA assay system showed good screening quality and reproducibility. Together with its higher throughput, rich real-time information, HTRA assay system could effectively provide biologically relevant cytotoxicity information for human health risk assessment.

**1972 Advancing Human Health Risk Assessment: Charting a Course through Committee Recommendations.**


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Over the last dozen years, many national and international expert groups have weighed in on specific improvements to risk assessment. Many of their stated recommendations are mutually supportive, but others appear conflicting, at least in an initial assessment. This effort synthesizes these opinions, identifies areas of consensus and sources of differences, and recommends a biological-centric, practical course forward for risk assessment, which includes: (1) Incorporating a clear problem formulation process at the outset of the assessment to ensure the level of complexity in design of the risk assessment is appropriate for informing the relevant risk management decision; (2) Use of mode of action (MOA) information and an understanding of the relevant biology as the key, central organizing principle for dose-response assessment; (3) Integrating MOA information into dose-response assessments using existing guidelines for noncancer and cancer assessments, and applying this knowledge with toxicokinetics to enable interpretation of human biomonitoring data in a risk context; (4) Using the tiered, iterative approach developed by the World Health Organization International Programme on Chemical Safety as a scientifically-sound framework for risk assessment of combined exposures (chemical mixtures). While scientifically-based defaults will remain important and useful when data on MOA or other data to refine an assessment are absent or insufficient, assessments should always strive to achieve the ultimate goal—to use 21st century knowledge of biological processes, dose-response, and chemical interactions at the molecular, cellular, organ and organism levels to minimize the need for extrapolation and reliance on default approaches.

**1973 Assessment on Health Risk and Risk Factors Related to Lead Exposure by Ingestion of Aquatic Animals from the Overflow Marsh.**

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The previous study identified that aquatic animals in Loeng Puay marsh were lead polluted. The lead concentrations in River snails were exceeded the food standard. This survey research aimed to determine health risk on ingestion of lead-contaminated aquatic animals from Loeng Puay marsh and the adverse health effects. The interviews of 75 residents at Loeng Puay on the consumptions of the aquatic animals and adverse health effects related to lead toxicity were conducted. The lead concentrations in aquatic animals (n=100) i.e. Nile tilapia, Common silver barb, River snail and Golden apple snail were analyzed by using Atomic Absorption Spectrophotometry. Health risk assessment was performed according to the U.S.EPA guideline (2004). The correlated factors with the adverse health effects were identified by the univariate analysis.

The highest lead concentration in all kind of aquatic animals was found in River snail and was used as represent concentration for health risk assessment on the acute effect. By considering the total intake rate of all aquatic animal consumption in the day, the maximum dose of lead exposure was 26.46 µg/day which was higher than the provisional tolerable weekly intake (PTWI=25 µg/kg). The adverse health effects related to lead toxicity were reported from 46.67% of residents. Most of symptoms were disorders of joints and skeleton, numbness, and acroesthesia, respectively. The univariate analysis identified that the fishermen had the highest risk on lead exposure (justified by adverse health effect) when compared to other occupations (OR=7.07, 95% CI: 2.20-23.81, p<0.001).

The findings indicated the health risk from high dose ingestion of lead-contaminated aquatic animals. However, the potential health risk depends on amount of food ingestion and personal factors. There should be continuously environmental monitoring of the lead accumulation in this overflow marsh for the safety of food chain and the health surveillance program.

**1974 Integrating Hazard Characterization Approaches for Evaluating the Potential of Consumer Products to Cause Asthma.**

I. Paterson1, A. Maiier1, M. J. Vincent and B. K. Gadagbui1. TERA, Cincinnati, OH.

Concerns have been raised regarding the potential for consumer products, including cleaning products to cause or exacerbate asthma or asthma-like responses. Although many forms of asthma are inflammation-based, some low-molecular-weight chemicals have been shown to trigger immunoglobulin E (IgE) independent occupational asthma. Single exposures to high concentrations of chemical irritants are also known to elicit an asthma-like response: reactive airways dysfunction syndrome (RADS). RADS can occur within hours of the initial exposure and may continue, as non-specific bronchial hyper-responsiveness, for extended durations. Exposure to irritants may be a trigger for respiratory symptoms in individuals with pre-existing asthma. Current methods cannot adequately assess the potential for consumer product ingredients to trigger asthma or asthma-like responses; epidemiological studies can only measure possible effects associated with a multitude of chemicals and products, and no single animal model can reliably replicate the complexity of an asthma-like response in humans. In order to characterize asthma and respiratory related hazards associated with consumer products, a decision system is needed that incorporates existing guidance, frameworks, and models. To develop such a tool, we compiled and evaluated in vivo, in vitro, and in silico methods that may provide data, or insight, to predict potential asthma or asthma-like responses (e.g., respiratory sensitization) and noted strengths and weaknesses associated with each method. We collaborated with asthma research experts to refine our findings and approach. Despite the wealth of information on asthma, current guidelines, biosays, and computer models cannot definitively identify whether a particular ingredient, or chemical, causes or exacerbates asthma or asthma-like responses. However, potential predictors of allergy-induced asthma, such as respiratory sensitization, are useful to assess the likelihood that a particular chemical, ingredient, or product may be associated with asthma induction.
and D.

EPA’s benchmark dose model software (BDMS) in accordance with standard EPA available studies that meet EPA’s study quality criteria and OSFs were derived using nalizes the PAH Mixtures guidance document. A literature search identified the (BaP) is due to be released for public review in 2013. The document will present an mediation could increase with the proposed changes in RPFs. Additional issues in-

ceedances of heath-based soil screening levels using current and proposed RPFs. Using EPA’s Records of Decision database for Superfund sites, we investigated the potential impacts of proposed changes in regulatory toxicology of PAHs on regulatory decision-making. Specifically, we investigated the degree to which changes in PAH RPFs could alter conclusions about health-based screening at DOD sites. Using EPA’s Records of Decision database for Superfund sites, we identified 11 DOD sites with 22 exposure units in which PAHs were identified as chemicals of concern in surface or subsurface soil. Site data were evaluated for ex-

cceedences of heath-based soil screening levels using current and proposed RPFs. Results indicate that the percentage of sites exceeding a screening level would in-
crease; up to twice as many sites for some individual PAHs (e.g., chrysene). In add-
dition, the magnitude of exceedance would increase for all sites. The results suggest that both the number of sites and the areal extent of individual sites requiring re-

mediation could increase with the proposed changes in RPFs. Additional issues in-
clude lack of background data for PAHs newly added to the risk assessment para-
digm and methods for assessing undetected PAHs. This work was supported in part by a grant from the Strategic Environmental Research and Development Program.

1976 Rapid Risk-Based Response to a Crude Oil Spill.

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Issues surrounding pipeline safety and the potential health and environmental im-

pacts that can result from pipeline accidents and malfunctions have led to consider-
able debate lately across North America. A number of recent high-profile oil spills in areas that are populated highlight the need for risk assessment to be completed rapidly to inform risk-based decision making. An example of such an assessment was completed in response to a break at an underwater pipeline crossing in which a sweet crude oil was released in a watercourse, upstream of several residences, active livestock farming operations, recreational properties and a municipal drinking water intake. Within 24-hours, a clean-up and contaminant sampling strategy was developed and implemented, and continued over several weeks. Preliminary evalu-

ation and interpretation of the contaminant data was completed to aid in commu-
nity consultation, to identify the need for further monitoring, and to provide risk-
based decision making strategies to local authorities managing drinking water and recreational use access. The conclusions of an initial risk assessment completed two weeks after the spill served to provide reassurance to regulators and the general pub-
lic regarding the safety of drinking water for residents and livestock, resulting in the re-

opening of some drinking water intakes in the impacted area. The assessment considered the nature and levels of the contaminants that were determined to be present in the water column and sediments, the results of toxicity studies for the various contaminants, and the opportunity for exposure of people and animals to the contaminants. Following the analysis of additional sampling results, a more comprehensive risk assessment was completed. The final results of the assessment revealed that the clean-up efforts had successfully mitigated risk to human health and the local ecosystem.

1977 Validation Oral Slope Factors for Benzo(a)pyrene Using Whole Mixture Results.

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EPA’s draft Integrated Risk Information System Assessment of Benzo(a)pyrene (BaP) is due to be released for public review in 2013. The document will present an Oral Slope Factor (OSF) that will replace the current value of 7.3 (mg/kg-day)-1. The OSF will be coupled with new Relative Potency Factors (RPFs) when EPA finalizes the PAH Mixtures guidance document. A literature search identified the available studies that meet EPA’s study quality criteria and OSFs were derived using EPA’s benchmark dose model software (BDMS) in accordance with standard EPA policy and guidance. The OSFs and the proposed RPFs have been validated against the results of rodent mixture studies to determine if the BaP OSF in conjunction with EPA’s proposed RPFs properly estimate PAH mixture effects. Benchmark dose modeling Culp et al. (1998) data results in two BaP OSFs with reasonable model fits: forestomach – 1.23 (mg/kg-day)-1 and esophagus – 0.19 (mg/kg-day)-1. Culp et al. (1998) also tested two coal tar samples in the same study. BaP-Toxic Equivalents were calculated for the coal tar mixtures using the EPA proposed RPFs. A nominal concentration of 100 ppm was assumed for unusual PAHs for which concentrations were unknown. When compared to the whole mixture OSFs for two separate coal tar mixtures, the current BaP OSF overestimates carcinogenic risk by 20 to 31-fold. The BaP OSF based on forestomach tumors in Culp et al. (1998) overestimates risk by 3 to 5-fold, whereas the BaP OSF based on esophagus tumors in the same study gives no overestimation. If the unusual PAHs with new RPFs are present in coal tar at concentrations significantly higher than the assumed 100 ppm, then the BaP OSF based on esophagus tumors will overestimate the coal tar tumorigenic risks observed in the whole mixture studies. This validation exercise demonstrates that the EPA proposed RPFs have to be implemented side-by-side with an OSF for BaP that is less than 1 (mg/kg-day)-1 to ensure that risks from real world PAH mixtures are not overestimated by the RPF risk assessment approach.

1978 Cytotoxicity Test for Hazardous Substances by Using the RTCATM High-Throughput Assay.

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Most current toxicity tests require the use of laboratory animals and involve expo-

sure by various routes and monitoring whether animals die or exhibit any clinical signs of toxicity. Increasing societal concerns about animal use have led to the de-

velopment of alternative in vitro test methods that can significantly reduce and re-

fine the use of mammalian for this purpose. The emerging in vitro cell based assays endeavor to provide information about the mode or mechanism of action that causes cytotoxicity and use such information to develop cell-based in vitro tests that can adequately model and predict in vivo toxicity. A non-invasive biophysical ap-

proach for the detection of cellular changes, the RTCATM instrument was devel-

oped for screening hazardous substances. In the present study, 6 chemicals (sodium arsenate, cadmium chloride, colchicine, verapamil, propranolol and mercury chlor-

ride) were applied to human renal adenocarcinoma cell line (ACHN) at 11 serial diluting working concentrations, respectively. The net cellular responses (the Cell Index) were recorded for 72 hours after adding chemicals. The real-time and unique time-concentration-dependent response curves (TCRC) were obtained. The experimental design composed two stages: initial screening test (range finder test) and definitive test (more defined test concentrations). The better TCRC pat-

terns were observed in the latter. The patterns were associated with some toxicity pathways or mode of action for these chemicals. The lethal concentrations could be calculated, which could be used to predict the starting lethal doses in vivo test. Based on these preliminary findings, other 8 substances in other 6 cell lines will be tested in the future. This work will facilitate cellomic based test development and achieve the replacement of animals in acute oral toxicity testing.

1979 Advancements in Arsenic Research Suggest a Dose-Dependent Transition Concentration for Cancer Endpoints.

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Recent in vitro studies have been conducted in both animal and human primary bladder cells to investigate the potential mode of action for bladder cancer follow-

ing exposure to arsenic compounds. Results from 24 hour in vitro gene expression studies with human uroepithelial cells treated in culture with mixtures of inorganic arsenic and its pentavalent or trivalent metabolites provide evidence of a common suite of gene changes consistently identified for a number of key signaling path-

ways: oxidative stress, protein folding, growth regulation, metallothionein regula-

tion, DNA damage sensing, thioredoxin regulation, and immune response. Lowest observed effect levels (LOELs) ranged from 0.6 – 6.0 μM total arsenic and no ob-
served effect levels (NOELs) ranged from 0.18 – 1.8 μM total arsenic. Benchmark dose modeling of the responses indicated lower confidence limits (BMDLs) ranging from 0.09 – 0.58 μM total arsenic for the eight genes most commonly significantly expressed across individual samples for the trivalent arsenical mixture, and from 0.35– 1.7 μM for total arsenic in the pentavalent arsenical mixture. These studies
provide the first evidence of no effect levels for arsenical-induced cell signaling perturbations in normal human cells exposed to biologically plausible concentrations of arsenic. Results from longer term exposures, again in primary human cells, provide consistent evidence that also suggest a dose-dependent transition for arsenic. Results of these in vitro studies, in combination with evidence from epidemiological studies, provide the basis for a shift in the approach for conducting a cancer risk assessment for arsenic. It also provides support to the conceptual demonstration that a dose-dependent transition in responses from those representing adaptive change to those that may be key events in the development of cancer endpoints.

1980 Refining the Application of a Database Uncertainty Factor in Human Health Risk Assessment.

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In human health risk assessment (HHRA), lack of information on a chemical is typically accounted for by the use of uncertainty factors (UFs). Typically, UFs are applied to account for five major areas of uncertainty: interspecies variation, intraspecies variation, extrapolation from subchronic to chronic exposure duration, extrapolation from a LOAEL to a NOAEL, and an incomplete database. To reduce uncertainty inherent in HHRA, methods are needed to replace or better assign values to these UFs. In this preliminary analysis, we analyzed all chemicals from the USEPA’s Toxicity Reference Database (ToxRefDB) with complete databases (i.e., general toxicity studies in two species, developmental toxicity studies in two species, and a multigeneration reproductive study) to assess whether a database UF (UDF) of 3 or 10 is sufficient to account for missing data. A subtractive approach was employed in which the points of departure (PODs) based on reported LOAELs and NOAELs in the ToxRefDB were compared for the complete database and following removal of selected studies. Removal of developmental toxicity studies from the database increased the POD by >3-fold in 59/186 (31.7%) chemicals and by >10-fold in only 1 chemical. Removal of multigeneration reproductive toxicity studies from the database increased the POD by >10-fold in 18/186 (9.7%) chemicals and by >10-fold in 4/186 (2.2%) chemicals. Removal of general toxicity studies from the database increased the POD by >3-fold in 59/186 (31.7%) chemicals and by >10-fold in 27/186 (14.5%) chemicals. This analysis will need to be supplemented with studies from a broader spectrum of chemical types (ToxRefDB is composed almost entirely of pesticides). When completed, this analysis could provide a method to systematically and empirically select the UFD in cases where chemical-specific information to inform selection of the UFD is lacking. The views expressed in this abstract are those of the authors and do not necessarily reflect the views or policies of the U.S. EPA.

1981 Short-Term Exposure to Perfluoroalkyl Acids Causes Increase of Hepatic Lipid and Triglyceride in Conjunction with Liver Hypertrophy.

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Persistent presence of perfluoroalkyl acids (PFAAs) in the environment is due to extensive use of industrial and consumer products. These chemicals activate peroxisome proliferator-activated receptor-alpha (PPARα) in liver and alter lipid metabolism. The current study was designed to evaluate liver toxicity of perfluorooctanoic acid (PFOA), perfluorolauronic acid (PFNA), perfluorohexane sulfonate (PFHXS), and perfluorophosphonic acid (PFPA), with emphasis on hepatocellular hypertrophy and steatosis. SV129 wild-type (WT) and PPARα-null (Null) adult male mice were dosed for 7 days with vehicle, PFOA, PFNA, PFHXS (10 mg/kg), and PFPA (20 mg/kg); and WY14643 (50 mg/kg) was a positive control. Mice were killed 24 hours after the last treatment. Liver samples were collected for biochemical analysis of triglyceride (TG) and DNA content. Frozen 6 μm sections of liver were stained with Oil Red O for lipid and used for morphometric analysis. Liver weights were elevated in both WT and Null mice in all of the treatment groups, except in Null mice of the PFPA and WY groups. Morphometric analysis revealed an increase in cell size in WT and Null livers exposed to PFOA, PFNA, PFHXS or PFPA, except for the PFPA and WY groups in Null mice. This pattern of change is consistent with the reduced DNA content per mg liver. In the Oil Red O stained sections, WT liver showed increased lipid accumulation in all treatment groups; whereas in Null livers, this was seen only after PFNA and PFHXS treatment. Similarly, elevated TG level was found in PFOA-exposed WT but not in WT-exposed mice, and increased TG was seen in Null mice only after PFNA treatment. Null livers had more lipid and TG than WT livers, both in control and treated mice. These results indicate that PFAAs induce liver hypertrophy and steatosis in WT and the involvement of PPARα is suggested by observations in Null mice. (This abstract does not necessarily reflect US EPA policy.)

1982 Structure-Activity Relationships for Perfluoroalkane-Induced Interference with Rat Liver Mitochondrial Respiration.

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Perfluorinated alkanes (PFAAs) represent a broad class of commercial products designed primarily for the coatings industry. Unfortunately, residues have been detected in a variety of species worldwide in association with cryotoxicity testing. Although some effects of PFAA exposures are linked to activation of the PPARα nuclear receptor, interference with mitochondrial bioenergetics has been implicated as an alternate and possibly important mechanism of cytotoxicity. The current investigation was designed to explore structure-activity relationships for the various modalities by which PFAAs interfere with mitochondrial respiration. Freshly isolated rat liver mitochondria were incubated with one of 16 different PFAAs. The effects of each PFAA on mitochondrial respiration was measured at five different concentrations and dose-response curves generated. All PFAAs tested inhibited mitochondrial respiratory control, mostly by stimulating state 4 and inhibiting state 3 respiration with the EC20 ranging from 1 μM to 413 μM. The perfluorooctyl sulfonamides were most potent with an EC20 between 1 and 11 μM. The results are consistent with evidence that the perfluorinated carboxylic acids stimulate the mitochondrial permeability transition whereas the sulfonamides act as protons uncouplers of oxidative phosphorylation. In all cases, there was a pronounced increase in potency with increasing carbon number, with a prominent inflection in potency between the six and eight carbon congeners. The results provide a foundation for classifying PFAAs according to specific modes of mitochondrial toxicity and establishing structure-activity evidence-based initial estimates of safety for PFAA congeners for which minimal in vivo toxicity testing currently exists. (This work was funded by NIEHS NTP contract 273200628005C).

1983 Determination of Polychlorinated Biphenyls (PCBs) and Hydroxylated Metabolites (OH-PCBs) in Human Blood Serum from Two Populations in East Chicago, IN and Columbus Junction, IA.

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PCBs are persistent and bioaccumulating toxic pollutants which pose health risk to humans and organisms. Although commercial production of these compounds was reduced and then banned in the 1970s, they are still present in our environment and found in human blood serum. In this study, we determined PCBs and their metabolites in human blood serum in these two populations. Human blood serum samples were collected as part of the Airborne Exposure to Semi-Volatile Organic Pollutants (AESOP) study from East Chicago, an industrialized area with known high PCB exposures in the past, and Columbus Junction, a rural area with no recognized historical PCB contamination. Our methods, which emphasize rigorous quality assurance and control (QA/QC), enabled us to evaluate all 209 PCB congeners and a sub-set of OH-PCBs congeners. After a series of extraction and clean-up procedures, samples were analyzed using gas chromatography with tandem mass spectrometry (GC-MS/MS) for PCBs and gas chromatography with electron capture detection (EC-D) for OH-PCBs. Our results show that the sum of PCB congeners ranged from non-detect to 658 ng/g.f.w. (median = 33.5 ng/g.f.w.) and the sum of the four major OH-PCB congeners ranged from non-detect to 1.2 ng/g.f.w. (median = 0.07 ng/g.f.w.). We conclude that PCBs and OH-PCBs are detected in human blood serum from populations living in East Chicago, IN and Columbus Junctions, IA.

1984 Associations among Polychlorinated Biphenyls (PCBs) and Chlorinated Pesticides and Serum Lipids in Residents of Anniston, Alabama.

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Associations among total cholesterol, triglycerides, and high density lipoprotein cholesterol (HDL) and serum polychlorinated biphenyls (PCBs) and chlorinated pesticides concentrations were examined in a sample of residents of Anniston, Alabama, participants of the Anniston Community Health Survey (ACHS).
For this analysis, nine PCBs with different number of chlorines (in brackets) and toxicological characteristics were selected: PCBs 74 (4), PCB 99 (5), PCB 118 (5), 153 (6), 170 (7), 187 (7), 196-203 (8), 206 (9), and 209 (10) were included. Four chlorinated pesticides were also included in the analyses: β-HCH, Oxychlordane, trans-Nonachlor, and p,p′-DDE.

Study includes 765 subjects with PCBs and lipid measurements. We wanted to minimize the effect of altered lipid levels due to presence or progression of disease(s) in studying the associations. Persons on lipid lowering medications, with dyslipidemia, diabetes, and coronary heart disease were excluded from the statistical analyses. Multiple linear regression models adjusted for age, race, gender, and BMI were used to analyze the data. Other major CHD risk factors such as smoking, exercise, and family history of heart disease were also examined. Only higher chlorinated PCBs 206 and 209 were related to total cholesterol and triglycerides; trans-Nonachlor was associated with triglycerides only. No association with PCBs or pesticides was found for HDL.

Interpretations of these results are limited. In particular, potential sequence of events relating lipid levels and PCB exposure is complex and uncertain and can only be elucidated with longitudinal study design.

1985 Persistent Organic Pollutants and Transthyretin-Bound Thyroxin in Plasma of Inuit Women of Childbearing Age.

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The Inuit population of Nunavik (Northern Quebec, Canada) is highly exposed to persistent organic pollutants (POPs) through their traditional diet which comprises fatty tissues from marine mammals. Some POPs – i.e. hydroxylated metabolites of polychlorinated biphenyls (PCBs), pentachlorophenol (PCP) and perfluorooctane sulfonate (PFOS) – are known to compete with thyroxin (T4) for binding sites on transthyretin (TTR), a T4 transport protein located in plasma and cerebrospinal fluid. Displacement of T4 from TTR could result in decreased T4 delivery to the developing fetus and in turn delayed growth and impaired neurodevelopment in infants. We set out to test the hypothesis that POPs or their metabolites decrease circulating concentrations of T4 bound to TTR (T4-TTR) in Inuit women of reproductive age. We measured T4-TTR concentrations in archived plasma samples obtained from 120 Inuit women (18-39 year old) by combining native polyacrylamide gel electrophoresis and liquid chromatography-tandem mass spectrometry (LC-MS/MS) techniques. Total T4 concentration was measured by LC-MS/MS, while those of TTR and thyroxin binding globulin (TBB) were determined by gel densitometry and an ELISA assay, respectively. POPs levels had been previously determined in those samples. The mean T4-TTR concentration was 8.4 nmol/L (SD = 2.4) with values ranging from 2.9 to 14.4 nmol/L, representing on average 6% of total T4 circulating concentration. Plasma levels of HO-PCBs, PCP or PFOS were not correlated to T4-TTR concentrations. Linear regression analysis revealed a weak correlation of TTR, TBB and total T4 concentrations were significant predictors of T4-TTR levels (adjusted R-square=0.26, p<0.001), but not POPs levels. Our results suggest that actual circulating levels of POPs in Inuit women of childbearing age are not high enough to affect TTR-mediated thyroid hormone transport.

1986 Effect of TCDD on Peripheral Hormones Regulating Feed Intake and Energy Balance in Rats.

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A lethal dose of 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) brings about a drastic body weight loss, accompanied by a decreased peripheral growth rate in rats. The decline of body weight primarily results from severe hypophagia but at present next to nothing is known about the factors underlying the reduced feed intake. The most important peripheral hunger signal in the body is ghrelin, mainly secreted into blood by stomach cells, whereas the principal satiety hormone is leptin produced by the white adipose tissue. Practically no data exist on the impact of TCDD on these hormones. In the present study, we exposed TCDD-sensitive Long-Evans (Turkil/AH; L-E) and TCDD-resistant Han/Wistar (Knöpfle/HW) rats to 100 μg/kg PCDD ig. and euthanized the rats at days 1, 4 or 10. The dose selected is lethal to all L-E rats but sublethal to H/W rats. A feed-restricted control (FCR) group of L-E rats was included in the study to help distinguish between primary and secondary effects. In addition to leptin and ghrelin, also glucagon was determined in serum samples with the Bio-Plex Suspension Array System. By 10 days, body weight dropped by about 30% in TCDD-treated and FRC L-E rats but only by some 5% in TCDD-treated H/W rats. Concurrently, serum leptin decreased to non-detectable levels in FRC. It diminished also in TCDD-treated L-E and H/W rats but less prominently. Serum ghrelin was elevated by TCDD in L-E rats at all time-points, and in H/W rats on day 10; in FRC, the increase was lesser on day 4 and comparable on day 10. Glucagon took an upward course in TCDD-treated rats reaching a nine-fold increase vs. control on day 10. These data reveal that the major peripheral signals of energy balance remain intact and change appropriately in lethally TCDD-treated L-E rats. However, they do not lead to the normal behavioural response (i.e. feeding) to rectify the tilted energy status.
PCBs, industrial chemicals and persistent environmental pollutants, are found in many rural and urban settings. Previous studies have shown that the treatment of rats with PCB126 causes a significant disruption of hepatic metal homeostasis that could potentially alter antioxidant defenses. The current study is focused on this metal disruption, in particular on Cu, Zn and Se. Two copper-containing proteins were investigated, tyrosinase (TYR) and cytochrome c oxidase (COX), along with metal chaperones ceruloplasmin (CP), sceloprotein p (SelP) and metallothionein (MT). These proteins cover a wide range of functions from intra- and extracellular metal trafficking (MT, SelP and CP), pigment production (TYR) and electron transport (COX). An animal study was conducted where 56 rats were fed one of three AIN diets containing levels of copper (2.6 & 10 ppm). After three weeks, animals were given a single IP injection of PCB126 (1 umol/kg or 5 umol/kg in corn oil) and euthanized two weeks later. The expression levels of these proteins were investigated by qRT-PCR, ELISA, and western blot. Metal analysis showed a decrease of Zn and Se, but an increase of Cu in the liver. Serum CP concentration and hepatic mRNA levels were increased with dietary copper and PCB treatment (although the latter not significantly). TYR was expressed in the liver both transcriptionally and translationally a new finding. COX was decreased with PCB126 exposure and had no statistical association with dietary copper. SelP was unchanged either by PCB126 or dietary copper, not reflecting the decrease of hepatic Se. MT was highly increased by PCB126 at the 1 umol/kg dose. The increase of metallothionein seen along with the higher binding affinity of Cu potentially explains the changes seen with Zn. Although metallothionein may contribute to the disruption of hepatic metal homeostasis, it fails to fully explain the Se and Cu changes.

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PCBs, a group of 209 individual congeners, were widely used as industrial chemicals and are ubiquitous and persistent human and environmental contaminants. Even though PCBs are carcinogens in rodents, they are still in use in closed applications. Lower chlorinated biphenyls can be metabolized to dihydroxy metabolites and further oxidized to quinoid metabolites both in vitro and in vivo. Quinoid metabolites may form adducts at nucleophilic sites of proteins like cytochrome c, an essential component in the mitochondrial electron transport chain. We hypothesized that the PCB-quinones covalently bind to cytochrome c thereby causing defects in mitochondrial function. LC-MS was used to detect adducts of cytochrome c with selected PCB-quinones in vitro. SDS PAGE gel electrophoresis followed by NBT staining was employed to separate the adducted proteins. Trypsin digestion and LC-tandem MS were applied to identify the amino acid binding sites on cytochrome c. In addition, cross-linking of cytochrome c was observed on the SDS PAGE gel. Different conditions, such as pH, incubation time and concentrations of PCB quinones, influence the formation of cross links. Lysine (K27, K39, K54/56, K73/74) and glutamate (E61, E62) were identified as binding sites by LC-tandem MS. Software simulation showed conformation changes of adducted cytochrome c. These preliminary data provide evidence that covalent binding of PCB quinone metabolites to cytochrome c may be included among the toxic effects of PCBs. (Supported by NIHES Superfund Program P42 ES013661.)

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Increased expression of brominated flame retardants and incineration of bromine-containing materials has lead to an increase in the presence of polybrominated dibenzo-p-dioxins and dibenzofurans (PBDD/Fs) in the environment. Measurable amounts of PBDD/Fs have been detected in sediment, seafood, and human serum. Previous studies indicated that an acute single exposure to 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD), 2,3,7,8-tetrabromodibenzofuran (TBBF), 2,3,7,8-tetrachlorodibenzo-furan (TCDF), 1,2,3,7,8-pentachlorodibenzofuran (1PeCDF), 1,2,3,7,8-pentachlorodibenzo-p-dioxin (1PeCDF), 2,3,4,7,8-pentachlorodibenzo-furan (4PeCDF), 2,3,4,7,8-pentabromodibenzofuran (4PeBDF), and 2,3-dibromo-7,8-dichlorodibenzofuran (DBDDCD) suppressed antigen-specific antibody forming cell (AFC) responses in B6C3F1/N mice, and that the brominated compounds were more potent than their chlorinated analogs. In addition, we evaluated expression of xenobiotic metabolizing enzyme (XME) and immune-relevant genes in the livers from these mice. Gene expression was measured using qRT-PCR. TCDD, TBBF, TCDF, 1PeBDF, 1PeCDF, 4PeBDF, 4PeCDF, and DBDDCD up-regulated the phase I XME genes Cyp1a1 and Cyp1a2. Based on estimated ED50, the rank order of potency for Cyp1a1 was TCDD>DBDDCD>4PeBDF>1PeBDF>1PeCDF>4PeCDF>1PeCDF. For Cyp1a2 was TCDD>TBBF>1PeBDF>4PeCDF>4PeBDF>TCDF>1PeCDF. TCDD, 4PeBDF, TBBF, TCDF, DBDDCD altered expression of phase II XME, phase III XME, and the thyroid transport protein transthyretin (Ttr) genes. 4PeCDF and 1PeBDF altered a phase II XME and Ttr genes. Additionally, 4PeBDF, TBBF, TCDF, and DBDDCD downregulated several genes that have been associated with antibiotic production and/or inflammation. Collectively these changes in gene expression are consistent with previously reported suppression of the AFC response and induction of aryl hydrocarbon-mediated and thyroid hormone responses. This abstract does not necessarily reflect the policies or views of NIH.
**1993 PBDE-100 Induces Mitochondrial and HepG2 Impairment.**

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INTRODUCTION: Polychlorinated diphenyl ethers (PBDEs) have been used in diverse products and are ubiquitous contaminants in sediments, in biota and are also found in human tissues, raising concerns about its toxicity. Many studies have reported liver toxicity and induction of apoptosis by mitochondrial dysfunction. OBJECTIVE: The aim of this work was to investigate the mechanisms of toxicity using HepG2 cells and isolated rat liver mitochondria. METHODS: Briefly, the effects of BDE-100 (0.1-50 μmol/L) was assessed on the mitochondrial respiration; Mitochondrial Membrane Potential (ΔΨ); mitochondrial swelling; interaction with membrane using 1-anilino-8-naphthalene sulfonate (ANS) and 1,6-diphenyl-1,3,5-hexatriene (DPh); Ca2+ release and mitochondrial ATP levels with the luciferin-luciferase system using isolated rat liver mitochondria. Furthermore, cytochrome c was investigated by testing cell viability with (4,5 dimetilthiazol-2-il) -Δψ and Sulphorhodamine B. RESULTS: In higher concentrations BDE-100 was able to induce mitochondrial alterations, and interact with the mitochondrial membrane, inhibiting the phosphorylation; leading to dissipation of the ΔΨ, deregulation of calcium homeostasis and mitochondrial swelling add a reduction in mitochondrial ATP content. In addition, it was observed inhibition the proliferation and reduce viability of the cells showing its cytotoxic action. CONCLUSIONS: These results suggest the formation of pores in the mitochondrial membrane and alteration in mitochondrial structure and function, leading to cell death due to cytotoxic effects in HepG2, perhaps through mitochondrial pathway. Word Keys: Mitochondria; Sulphorhodamine B. Supported by: FAPESP

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**1994 Genomic Plasticity and Polychlorinated Biphenyls: Telomerase Reactivation and AhR Desensitization in Human Keratinocytes after Long-Term Exposure to PCB126.**

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Polychlorinated biphenyls (PCBs) are environmental pollutants and dioxin-like PCB126 (3,3',4,4', 5-pentachlorobiphenyl) is classified by IARC as a human carcinogen. Chromosome ends have 'telomeres' which shorten with age in somatic cells. Lengthening of telomeres and high telomerase activity, the telomere maintaining enzyme, are key steps in carcinogenesis. To examine if PCB126 carcinogenicity is mediated through this mechanism, we exposed immortalized human skin keratinocytes (HaCat) for 90 days to 5 μM PCB126. Every sixth day cells were re-seeded to telomerase activity, telomere length, C-Myc, hTERT and hTR (telomerase components), TRF1 and TRF2 (reduce telomerase access to telomeres) mRNA. CYP1A1 mRNA and activity, superoxide and H202 levels were determined. PCB126 caused an increase in CYP1A1 mRNA and activity, TRF1/2 mRNA, and superoxide and H202 and a reduction of telomerase activity (50%), hTERT and hTR mRNA (10%), telomere length (40%), and cell growth from Days 6 to 48. From Day 94 on, an increase in cell growth, C-Myc, hTERT, and hTR mRNA levels (to 130%), reactivation of telomerase activity (to 100%), elongation of telomere length (to 90%), and a decrease in TRF1 and TRF2 mRNA were observed. In addition, from Day 78 PCB126 no longer activated the AhR response (CYP enzymes) and no mutation were found on the AhR ligand binding region. Microarray results confirmed an increase in expression of cell growth genes on Day 54 and desensitization of AhR-response on Day 78. This study shows for the first time a telomerase reactivation, telomere lengthening, and increased cell growth after telomeres were significantly shortened by PCB-exposure in human cells. Possible mechanisms include de novo C-Myc amplification (telomerase) and/or selection of a small subpopulation of cells. Either way, this work adds a new toxicity pathway to PCBs, and a plausible mechanism of carcinogenic PCBs in related contaminants to be considered in their safety evaluation and risk assessment.

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**1995 Exposure to PCB 126 Triggers Antioxidant Defense through Cross-Talk of Caveolae and Nrfl2 Signaling.**

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Environmental toxicants such as polychlorinated biphenyls have been implicated in the promotion of multiple inflammatory diseases including cardiovascular disease, but information regarding mechanisms of toxicity and cross-talk between relevant signaling pathways is lacking. We have reported that coplanar PCBs promote endothelial cell activation through the lipid microdomain caveolae, and the loss of caveolin-1 (Cav-1) ameliorates these detrimental effects. We have also shown that PCB 126 can activate the antioxidant transcription factor Nrfl2 resulting in upregulation of antioxidant genes and downregulation of inflammatory markers. Normally, Nrfl2 is sequestered in the cytoplasm and degraded through inhibitory action of Keap1, but upon activation via toxicants such as PCBs, can enter the nucleus and activate the transcription of a battery of protective genes. Our previous data suggests downregulation of Cav-1 and upregulation of Nrfl2 protects against PCB-induced cellular dysfunction, but here we show for the first time an example of cross-talk between these two cellular signaling pathways. To examine the cross-talk between Cav-1 and Nrfl2 pathways in PCB-induced inflammation we silenced Cav-1 in vascular cells. Importantly, Cav-1 silenced cells treated with PCB126 resulted in increased levels of Nrfl2-ARE binding determined by EMSA. Also, in cells treated with PCB 126, silencing of Cav-1 resulted in decreased protein levels of both inhibitory Keap1 and Fyn kinase, which both have been previously to be implicated in Nrfl2 deactivation. We also show that Keap1 levels were significantly decreased in livers from Cav-1 KO mice when compared to control C57Bl6 mice. Finally, Cav-1 silencing allowed for a more effective antioxidant response, as observed by higher levels of the antioxidant genes glutathione s-transferase (GST) and NADPH dehydrogenase quinone 1 (NQO1) in cells exposed to PCB 126. Ultimately, these data introduce novel cross-talk between Cav-1 and Nrfl2 and implicates the ablation of Cav-1 as a protective mechanism of PCB-induced cellular dysfunction and inflammation.

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**1996 Fibroblast Growth Factor 21 (FGF21) is a Target Gene of the Aryl Hydrocarbon Receptor (AhR).**

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Dioxins, such as 2, 3, 7, 8-tetrachlorodibenzo-p-dioxin (TCDD), are environmental pollutants. The toxic effects of TCDD are well documented mainly through activating the aryl hydrocarbon receptor (AhR). However, the underlying mechanisms for TCDD’s adverse effects, such as the wasting syndrome, are not well understood. Fibroblast growth factor (FGF) 21 plays critical roles in metabolic adaptation to fasting by increasing lipid oxidation and ketogenesis in the liver. The present study was performed to determine whether activation of the aryl hydrocarbon receptor (AhR) induces Fgf21 expression. In mouse liver, TCDD increased Fgf21 mRNA in both a dose- and time-dependent manner. Moreover, TCDD increased Fgf21 mRNA levels in livers of wild-type mice, but not of AhR-null mice. Chromatin immunoprecipitation assays indicate that TCDD increased AhR protein binding to the Fgf21 promoter (-105/+1 base pair). Diethylylphthalate (DEHP) decreased Fgf21 mRNA expression and DEHP pretreatment attenuated TCDD-induced Fgf21 expression, which may explain a previous report that DEHP pretreatment decreases TCDD-induced toxicities. In mouse liver, TCDD increased the mRNA of fatty acid uptake protein CD36 but decreased mRNA of de novo fatty acid biosynthesis enzyme fatty acid synthase (FAS). In addition to these findings in liver, TCDD induced Fgf21 mRNA in mouse white adipose tissue. In conclusion, TCDD induces Fgf21 expression via activation of the AhR-signaling pathway. Pharmacological manipulation of Fgf21 expression by AhR activators may provide an effective strategy for treating obesity, diabetes, and the metabolic syndrome. (Supported by NIH grants DK-081461, ES-09649, ES-019487, and RR-021940.)

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**1997 Human Transcription Factor Activation by Polychlorinated Biphenyls and Organochlorine Pesticides.**

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Introduction: Polychlorinated Biphenyls (PCBs) are persistent environmental toxins, present in 100% of the US adult population and theoretically predicted to act as ligands for transcription factors involved in xenobiotic/endobiotic detoxification, obesity and steatosis. These human receptors include pregnane xenobiotic receptor (PXR), arylhydrocarbon receptor (AhR) and possibly the liver-X-receptor-α (LXR). This study evaluates a PCB mixture, Aroclor 1260, and selected individual PCB congeners, as potential ligands for these receptors. Additionally, we also look at PXR activation by selected organochlorine pesticides. Methods: MTTR assay was performed to determine acute toxicity concentrations for Aroclor 1260. HepG2 cells were transfected with plasmids expressing human LXR and human PXR, and receptor-responsive plasmids including tk-LXR-RE-luciferase, pGL3-PXR-RE-luciferase or pGL3-AhR-RE-luciferase. Transfected cells were treated with ligands for LXR (T0901317:100nM), PXR (Rifampicin:10μM) and AhR

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**1998 SOT 2013 Annual Meeting 425**
Validation of QuEChERS-Based Extraction for the Determination of Organochlorine Pesticides in Liver and Serum Using GC-MS/MS.

Organochlorine (OC) pesticides are chemically stable, have low volatility, and low rates of degradation which lead to their persistence in the environment and classification as persistent organic pollutants (POP). OC pesticides bioaccumulate and biomagnify to greater extents in animals at higher trophic levels due to their high lipid solubility. These compounds generally affect the nervous system of the target organism. The mechanism of action involves the disruption of chemical ion movement in neurons but the major pathologic changes are observed in the liver and reproductive systems. Michigan State University Diagnostic Center for Population & Animal Health, Michigan State University, East Lansing, MI; 2Pathology and Diagnostic Investigation, Michigan State University, East Lansing, MI.

Potential human health risks associated with exposure to dioxin-like compounds (DLCs) are evaluated using toxic equivalency factors (TEFs). TEFs are single point estimates even though they are based on relative estimates of potency (REPs) that often span several orders of magnitude. One potential improvement to the TEF methodology would be to use the full distributions of REP values for each congener. WH0 recognized the value of such an approach during their most recent review of the TEF methodology but expressed concern that all REP values are not of equivalent quality or relevance. As such, we previously established a consensus-based framework that weights REPs based on REP quality and relevance and to develop a numerical approach for quantitatively weighting each REP using machine learning. Since that time, we have applied the quantitative weighting framework to the REP database to develop a numerical weight for each REP value, which in turn has been used to develop a weighted distribution of REPs for each congener, and have also developed 95% confidence intervals (CIs) around the percentile associated with the TEF values using the R statistical programming environment. The weighted distributions were generally tighter than were the unweighted distributions. Additionally, when examining where the current WHO TEFs fell on the weighted distributions for each congener, we found a lack of consistency across congeners, with values percentiles ranging from the 1st–99th percentiles. The CI for many of the congeners were also quite broad. For example, the TEF for 12378 PdCDF fell on the 47th percentile of the weighted distribution, but the CI ranged from the 29th to the 68th percentile. These calculations improve characterization of the variability and uncertainty inherent in the health risk estimates for this class of compounds, demonstrating that reliance on the point estimate TEFs could significantly under- or over-estimate risk. (This abstract does not reflect the policies or views of NIEHS or NCI.)

Phenylenyl isothiocyanate and Dithiocarbamate Restore Proangiogenic Properties of Cyplb1-Deficient Vascular Cells through Decreased NF-kB Activity and Oxidative Stress.

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Purpose: Cytochrome P450 1B1 (Cyp1B1) is expressed in the vasculature and loss of its expression/activity is associated with increased oxidative stress, increased thrombospondin-2, and NF-kB activation. This is concomitant with attenuation of angiogenesis in vivo and in vitro. Both phenylethyl isothiocyanate (PEITC), found in cruciferous vegetables, and pyrrolidine dithiocarbamate (PDTC) inhibit NF-kB activity. The objective of this study was to determine if inhibition of NF-kB by PEITC or PDTC restores the reductive and proangiogenic properties of Cyp1b1-deficient vascular cells.

Methods: Primary cultures of retinal endothelial (EC) and pericytes (PC) were prepared from Cyp1b1+/+ and Cyp1b1−/− mice. Cyp1b1−/− retinal vascular cells were incubated with either 1 μM PEITC or 10 nM PDTC for 24 h, unless indicated otherwise. NF-kB expression and activity were determined by Western blot analysis and luciferase reporter activity. Immunofluorescence staining was performed to visualize p65 localization. Oxidative stress was measured using dihydroethidium staining. The ability of Cyp1b1+/+ and Cyp1b1−/− vascular cells to undergo capillary morphogenesis in Matrigel was also determined after 8 h incubation with PDTC or PEITC. Rates of cell migration were compared using a tranwell assay. Results: PEITC and PDTC both inhibited NFKB p65 expression and activity. NF-kB inhibition decreased p65 staining and nuclear localization in Cyp1b1−/− retinal vascular cells. Inhibition of NF-kB restored capillary morphogenesis and migration of retinal vascular cells. Oxidative stress in Cyp1b1−/− retinal vascular cells was relieved with incubation with PEITC and PDTC.

Conclusion: Cyp1b1 expression/activity is essential for maintaining the normal proliferative, migratory, and reductive state of the vascular cells, and its alteration has significant impact on vascular development and angiogenesis.
2002 Cyp1b1-Deficiency Alters Structure and Function of Trabecular Meshwork via Increased Oxidative Stress.

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Cytochrome P450 1b1 (Cyp1b1) is a member of the cytochrome P450 superfamily of enzymes with mono-oxygenase activity. Although mutations in Cyp1b1 gene have been reported in patients with congenital glaucoma, the role Cyp1b1 plays in the development and function of trabecular meshwork (TM) remains unknown. Here we determined the impact of Cyp1b1 deficiency on the development and function of TM tissue in C57Bl6 mice. The intraocular pressures (IOPs) of Cyp1b1+/+/, Cyp1b1+/− and Cyp1b1−/− mice were measured non-invasively with a commercially available tonometer. The integrity of TM tissue of 1-week to 8-month-old Cyp1b1+/+, Cyp1b1+/− and Cyp1b1−/− mice were assessed by electron microscopy (EM). Oxidative stress in the TM tissue was evaluated by immunostaining for 4-hydroxy-2-sonoal (HNE), N-Acetylcysteine (NAC; 1.5 mg/g body weight, once every three days for three weeks, IP) was administered in 3-day-old Cyp1b1+/+ mice and the morphology of TM was assessed by EM. Our results showed a modest but significant increase in diurnal IOP of Cyp1b1−/− mice at 6-12 weeks of age. The 2-week-old Cyp1b1−/− mice presented ultra structural irregular collagen distribution in the TM tissue, which became more severe as mice aged. Increased HNE staining was observed in TM tissue of Cyp1b1−/− mice in vivo. Administration of NAC prevented the postnatal formation of lesions in Cyp1b1−/− mice. These results are consistent with our previous in vitro findings that TM cells prepared from Cyp1b1−/− mice exhibit increased oxidative stress and cellular defects, which are reversed when incubated with NAC. Thus, the metabolic activity of Cyp1b1 contributes to oxidative homeostasis and integrity of ultra-structure and function of TM.

2003 Macrophage Toxicity in Response to Particles Collected from Indium-Tin Oxide Production.

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Occupational exposure to indium compounds has recently been associated with lung disease among workers in the indium-tin oxide (ITO) industry. Previous studies have suggested that autoantibodies and reactive oxygen species (ROS) may play a role in the development of pulmonary lesions following indium compound exposure. However, the molecular mechanisms behind indium compounds' toxicity remain largely unknown. Thus, we aimed to uncover how compounds encountered in occupational health issue and as a result of exposure to indium, contribute to lung pathology. Therefore, various imaging techniques are being undertaken to evaluate the effects of the collected indium compounds on intracellular ROS production, lipid peroxidation, DNA damage, and cell proliferation over the same time course. These findings will provide a foundation for the molecular basis behind an emerging occupational health issue and as a result of exposure to indium, contribute to lung pathology. Therefore, various imaging techniques are being undertaken to evaluate the effects of the collected indium compounds on intracellular ROS production, lipid peroxidation, DNA damage, and cell proliferation over the same time course. These findings will provide a foundation for the molecular basis behind an emerging occupational health issue and as a result of exposure to indium, contribute to lung pathology.

2004 Role of Cytochrome P450 (CYP)1A Enzymes in Sex-Specific Differences in Hypoxic Lung Injury.

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Preterm male infants have a higher incidence of chronic lung disease (CLD) compared to females and hyperoxia contributes to its pathogenesis. Cytochrome P450(CYP) enzymes contribute to hyperoxia lung injury and the role of CYP1A enzymes remain largely unknown. We tested the hypothesis that mice will display sex-specific differences in hyperoxic lung injury and that this phenomenon will be altered in mice lacking the genes for CYP1A1 or 1A2 (male [M] and female [F] 8-10 wk old, wild type [WT], Cyp1a1-null, and Cyp1a2-null mice were exposed to hyperoxia [FO2=0.95]. Lung injury was estimated by lung weight/body weight (LW/BW) ratios, histopathology and immunohistochemistry for quantitation of neptanol infiltration. Levels of 8-iso PGF2α (lung) were determined by LC-MS/MS. Macrophages within 5 minutes. We also hypothesize that indium compound particles collected from Indium-Tin Oxide Production. (LW/BW) ratios, histopathology and immunohistochemistry for quantitation of neptanol infiltration. Levels of 8-iso PGF2α (lung) were determined by LC-MS/MS. Macrophages within 5 minutes. We also hypothesize that indium compound particles collected from Indium-Tin Oxide Production.

2005 Towards Elucidating the Pathophysiological Role of Reactive γ-Ketoaldehydes Formed through the Isoprostane Pathway of Lipid Peroxidation.

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Numerous environmental toxins cause oxidative damage to lipids, proteins, and DNA. Lipid peroxidation leads to the formation of reactive aldehydes amongst which the γ-ketoaldehydes formed via the isoprostane pathway of lipid peroxidation, termed isoketals (IsoKs), are the most reactive and injurious. Selective scavengers for IsoKs have been developed which have shown remarkable protection against oxidative damage in animal models of oxidative stress. However, the precise molecular processes preserved due to scavenging IsoKs have not been defined. This can, however, be elucidated studying the highly tractable organism, Caenorhabditis elegans (C. elegans). Accordingly, to identify the pathophysiological roles of IsoKs in oxidative injury, C. elegans are exposed to oxidative insult in the absence and presence of salicylamine. A fluorescent C. elegans strain containing a GFP transcriptional reporter for the SKN-1 target gene gst-4 (Pgst-4::GFP) was utilized for detection of altered gene expression in order to determine processes protected by reactive γ-ketoaldehyde scavengers. Late-stage L4 worms were dosed with a submaximal dose of a SKN-1 activating xenobiotic, juglone, for one hour and plated for recovery for one hour. Current studies show a nearly 4-fold increase in fluorescent intensity after a submaximal (LD50) dose of juglone (p-value = 0.02). These results demonstrate that a brief exposure to the xenobiotic juglone is sufficient in activating SKN-1 in C. elegans, and can be used in oxidative and xenobiotic stress studies. (Supported by T32EY007028-38 (T.I.N.)).

2006 High-Content Imaging Assay to Detect Drug-Induced Reactive Oxygen Species Generation.

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The disruption of cellular redox circuits can lead to an increase in reactive oxygen species (ROS) within cells, causing oxidative stress and eventually cell damage. Mitochondria are the major producers of ROS in the form of superoxide anion. Two of the electron transport chain complexes, Complex I and Complex III, are thought to be responsible for most of the ROS generated in mitochondria. Several drugs associated with oxidative stress have been shown to contribute to toxicity in the liver, heart, kidney and central nervous system. Thus, it is important to be able to screen new chemical entities that may cause an increase in ROS production in order to reduce compound attrition at the early drug discovery stage. Hence, we developed a 384-well format high content imaging assay that measures superoxide production. The effect of 60 compounds including thiazolidinediones, antipsychotics, antidepressants and anticonvulsants was assessed in this assay in transformed human liver epithelial cells (THLE) using the fluorescent dyes, Dihydroethidium (for superoxide levels) and Hoechst (for cell number). In a separate 384-well format high content imaging assay, we tested the effect of the compounds on the mitochondrial membrane potential (MMP) of the cells using the

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fluorescence dye, TMRM. Quantitative image analysis showed that the compounds could be grouped into four categories: (a) those that caused superoxide formation and a loss in MMP at similar concentration ranges (for example, astemizole, sorafenib), (b) those that caused superoxide production but had no effect on MMP (dasatinib), (c) those that caused a loss in MMP but had no effect on superoxide formation (tamoxifen, sertindole), and (d) those that had no effect either on superoxide formation or on the MMP over 24 hours (pioglitazone, riposetnone). Both high content imaging assays were robust and rapid and can be implemented within a screening paradigm to identify compounds that modulate oxidative stress and mitochondrial membrane potential.

2007 The Antioxidant Lipoic Acid Exacerbates Paraquat-Induced Cytotoxicity.

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In several countries the use of herbicides has become important in the preservation of sustainable agriculture. A widely used herbicide for broadleaved weed control is paraquat (PQ) known to be toxic to humans and animals. The treatment against PQ poisoning remains supportive with no antidotes or specific treatments available. Recognizing that PQ induces its toxicity primarily via oxidative stress-mediated mechanisms, modulating the levels of cellular antioxidants seems to serve as a potential therapeutic strategy. We studied the in vitro effects of the thiol-containing antioxidant lipoic acid (LA) in a human lung adenocarcinoma epithelial cell line. Incubation of control A549 cells with PQ resulted in time- and concentration-dependent increases in intracellular PQ levels with concomitant decreases in cell viability and mitochondrial membrane permeability (MMP) and increases in intracellular calcium concentrations ([Ca2+]i). Pretreatment of cells with LA did not cause any changes in any of the biochemical parameters measured with the exception of the MMP being significantly decreased. Co-treatment of A549 cells with LA and PQ potentiated the PQ-induced cytotoxicity as evidenced by the further exacerbation of PQ-induced decreases in MMP and increases in DNA fragmentation and [Ca2+]i. Chromatographic analysis (GC/MS/MS) showed that LA was primarily associated with cell membranes. These data suggest that LA does not offer any protective effects against PQ-induced toxicity. The mechanism(s) for its ability to modulate cell survival/death by modulating the cellular redox-regulated signal transduction in PQ-challenged cells is under investigation. This research project was supported by Natural Sciences and Engineering Research Council of Canada (NSERC).

2008 Lipid Droplets with Oxidized Fatty Acids and Triglycerides in Dendritic Cells: Possible Role in Antigen Presentation in Cancer.

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Immunosurveillance plays a critical role in control of tumor progression whereby dendritic cells (DC) are the most potent antigen presenting cells responsible for the development of immune responses. Our previous work has identified lipid droplets as potent regulators of DC's immune functions. Notably, DCs isolated from tumor-bearing mice or treated with tumor explant supernatants (TES) accumulated lipid droplets containing considerable amounts of PUFA (C18:2, C18:3, C20:4 and C22:6). Among those, the amounts of C18:2 were significantly higher compared to other PUFA species. Since accumulation of lipid droplets in DCs in cancer was mediated via Msr1 receptor, which is primarily responsible for the uptake of oxidatively modified lipids, we performed analysis of oxidized lipids in DCs using LC-ESI-MS. To investigate possible role of oxidized fatty acid in antigen presentation we used a model system with naive mice were grown in the presence of C18:2. Treatment of DCs with C18:2 in combination with a hydrophobic free radical generator, an azo-initiator AMVN, - but not with C18:2 alone - resulted in accumulation of oxidized lipid droplets and caused significant decrease in antigen presentation. Oxidated FFA containing one, two and three oxgens as well as oxidated triglycerycerols, TAGs, including truncated TAGs with m/z 766 [M-NH4]+, containing acyl corresponding to 9-oxo-nonanoic acid, were observed in DC grown in the presence of TES and DC treated with C18:2 and AMVN. Given that lipid droplets can directly co-localize with cellular compartments involved in antigen processing and formation of phagosomes, it is likely that accumulation of oxidated FFAs and TAGs in DC may be responsible for the loss of their immune-regulatory functions in cancer. Supported by grant with CA165065, NIOSH OH088282, NIH U19 AI068021, HL 70755.

2009 Mitochondrial Cardiolipin As a Substrate for Cytochrome C-Catalyzed Production of Oxygenated Lipid Mediators.

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Lipid mediators – central to the normal homeostasis and responses to stress and disease - are generated through oxygenation of polyunsaturated fatty acids (PUFA), such as linoleic acid (LA), arachidonic acid (AA) and docosahexaenoic acid. Their regulatory effects are believed to depend on a fine balance between PUFA esterification/recylation of phospholipids and on their hydrolysis by phospholipases A (PLA). We suggested that mitochondrial cardiolipins (CLs) can be a source of bioactive lipid mediators generated with the catalytic participation of cytochrome c (cyt c). In this study we employed models of rat traumatic brain injury (TBI) and mouse total body irradiation (TIR). Using oxidative lipidomics approach and MS/MS analysis, we found that TBI resulted in oxidation of polyunsaturated molecular species of CL and accumulation of its hydrolysis products such as oxygenated LA, AA and monoylo-CL. Similar, a significant increase of oxidized CLs in small intestine of TIR mice (9.5 Gy) was accompanied by accumulation of CL hydrolysis products. To generate oxygenated CL in vitro we utilized brain CL with its highly diversified polyunsaturated molecular species, and cyt c. We found that incubation of brain CL with cyt c in the presence of H2O2 yields a rich assortment of oxygenated CL species, hydrolysis of which by PLAs A1 and A2 generated multiple oxygenated fatty acids similar to those that were formed in vivo in brain after TBI and small intestine after TIR. An oxidation-specific lipoprotein lipase A2, was able to utilize peroxidized tetranolinoe-CL to yield different oxygenated species of linoleic acid and lyso-CLs. Thus, mitochondrial CL/cyt c represents a novel mechanisms involved in lipid mediators-generating pathways. Supported by NIH ES020693, ES021168, U19 AI068021, HL 70755; NIOSH OH008822, NS07651, N506187.

2010 Development of a Mitochondria-Targeted Nano-Complex of Imidazole-Substituted Oleic Acid As a Radiomitigator. A. Star1, 2, A. Kattalor1, 2, A. Amoscato2, 3, V. Tyurin2, 3, W. Seo1, 3, M. Epperly3, 4, J. Greenberger4, V. Tyurina2, 3, V. Kagan1, 2, 3, 4. Department of Chemistry, University of Pittsburgh, Pittsburgh, PA; 2Department of Environmental and Occupational Health, University of Pittsburgh, Pittsburgh, PA; 3Center for Medical Countermeasures against Radiation, University of Pittsburgh, Pittsburgh, PA; 4Radiation Oncology, University of Pittsburgh, Pittsburgh, PA.

Increasing likelihood of intended or accidental exposure to ionizing radiation dictates the necessity to develop effective medical countermeasures of radiation injury as has been recognized as a high priority both in the US and worldwide. No effective medical radiation countermeasures exist for intervention in acute and delayed radiation injuries are currently known. Based on newly discovered mechanisms of radiation damage – oxidation of cardiolipin by cytochrome c in mitochondria as a required stage in radiation-induced apoptosis - we designed and synthesized mitochondria-targeted triphenylphosphonium-conjugated imidazole-substituted oleic (TPP-OA) which prevented/mitigated cell death induced by irradiation and protected C57BL6 mice from total body irradiation. To improve therapeutic efficiency of TPP-OA we chose to employ branched polyethylene glycol (PEG) functionalized single walled carbon nanotubes (SWCNT) and use it as a carrier to deliver mitochondria-targeted TPP-OA to tissues. We found that loading of PEG-SWCNT with TPP-OA caused a marked extension of the life-span of TPP-OA in circulation. Moreover we showed that TPP-OA-nano-complex was more effective as radiomitigator than free TPP-OA. While the dose of TPP-OA in TPP-OA-nano-complexes was two times lower compared with free TPP-OA the mitigating effect of TPP-OA-nano-complexes was higher than that of TPP-OA alone. Importantly, we were able to deliver TPP-OA nano-complexes in radiosensitive tissue such as small intestine. These data warrant further studies aimed at the development of radioprotectors/radiomitigators with broad spectrum of applications in biomedicine and biodefense. Supported by NIOSH OH088282; NIH U19 AI068021, HL 70755, ES019304.

2011 XBP1, SYNV1 and Nrfr2 At the Crossroads of ER Stress and Oxidative Stress. T. Wu and D. D. Zhang. College of Pharmacy, University of Arizona, Tucson, AZ.

Transcription Factor Nrfr2 has long time been revealed as the master regulator of intracellular redox homeostasis. As an adaptive response to oxidative stress, Nrfr2 activates transcription of a battery of genes encoding antioxidant protein, detoxification
enzymes and xenobiotic transporters by binding to the cis-antioxidant response ele-
ments in the promoter region of these genes. Previous works by our lab and others
demonstrated that Nrf2 is subject to poly-ubiquitin-mediated proteasomal degra-
dation in a Keap1-dependent manner. Here we report that active form of XBP1, XBP1s suppresses Nrf2 and its downstream signal under ER stress condition,
through activation transcription of synovial apoptosis inhibitor 1 (SYVN1). SYVN1, also known as Hrd1, is an ER-associated degradation (ERAD) ubiquitin
ligase. We have determined that SYVN1 directly interacts with the Neh4 and Neh5
domain within Nrf2. Overexpression of SYVN1 attenuates Nrf2 signaling, whereas
knockdown of SYVN1 enhances expression of Nrf2 and its downstream genes. Fur-
thermore, SYVN1 accelerates the clearance of Nrf2 protein through promoting
ubiquitination of Nrf2. These findings demonstrate that XBP1 and SYVN1 are invol-
volved in regulating the Nrf2 pathway in a new Keap1-independent mechanism.
Moreover, our data revealed a possible crosstalk between ER stress pathway and ox-
idative responses via UPR and Nrf2 signaling pathway in order to protect cells
against environmental stress.

**2013 Generation of Reactive Oxygen Species by Process Materials from Indium-Tin Oxide Production.**

N. R. Fix1, K. M. Dunnik1, M. A. Badding1, A. B. Stefanik3, K. J. Cummings2, V. Castranova2 and S. S. Leonard1. 1Health Effects Laboratory Division, National Institute of Occupational Safety and Health (NIOSH)/CDC, Morgantown, WV; 2Division of Respiratory Disease Studies, National Institute of Occupational Safety and Health (NIOSH)/CDC, Morgantown, WV. The transition metal indium has been used for decades for various applications in-
cluding electronics, fusible alloys, and solar cells. Indium compounds usage has in-
creased dramatically based on the rise in demand of touch screens and flat panel
displays (LCDs). With this growth of industry, there is potential for increase of in-
dium lung disease among workers who produce, use, or reclaim indium-tin oxide (ITO). Inhalation exposure of indium samples can occur during various times of manufac-
turing. Materials from different sources, Excreta were collected from an ITO
production facility. While the pathogenesis of indium lung disease is unknown,
previous work has suggested a role for reactive oxygen species (ROS). Chemical
characteristics of the process materials will aid in determination of reactivity differ-
ences between compounds. Electron spin resonance (ESR), a common tool used for
measuring ROS, was used in both acellular and cellular exposures. Acellular sam-
ple were evaluated by combining 10 mg/mL process material, phosphate buffered
saline (PBS), 10mM hydrogen peroxide (H2O2), and 100mM DMPO (spin trap).
RAW 264.7 mouse macrophages, DMPO, and the same concentration
of composite were used in the cellular samples conducted in ESR. Scavengers and
chelators were used to define radical mechanisms. Results indicated that ventilation
dust (VD), tin-oxide (SnO2), and unsintered ITO (UITO) cause a greater increase in
ROS production than the other process material. H2O2 and O3 consumption
measurements were used to determine the source of the ROS. ESR studies com-
pared with measurement of ROS production will help to determine the mechanisms
behind indium lung disease. Data from this study will be used to determine possi-
ble hazards in occupational exposure of indium process material while increasing
the understanding of indium lung disease.

**2014 Lower Expression of Nrf2 Promotes Proliferation, Migration and Invasion of Prostate Cancer Cells.**

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The nuclear factor Nrf2 is known to play a critical role in cellular protection against
oxidative stress and cellular transformation. However, unabated nuclear accumu-
lation of Nrf2 is also known to reduce apoptosis, promote cancer cell survival and
drug resistance. Mutations in Infr2 and Nfr2 leading to nuclear accumulation of
Nrf2 in many cancers including prostate and breast cancer are known. These also
raise interesting questions regarding the role of Nrf2 in cancer metastasis and/or
metastasis progression that remains elusive. In this study we have investigated the
hypothetical role of loss of Nrf2 is associated with metastasis/metastasis progression.
We used less metastatic LNCaP and highly metastatic LNCaP derived C4-B2 prostate
cancer cell lines to test our hypothesis. The analysis revealed that highly metastatic
C4-B2 cells expressed higher Infr2 and lower Nfr2 levels as compared to less
metastatic LNCaP cells. We used control, Infr2 and Nfr2 shRNA to generate C4-B2
and LNCaP derived cells with altered levels of Infr2 and Nfr2 to determine the
role of Nfr2 in metastasis and metastasis progression in Soft agar colony formation
and X-CELLigence proliferation, migration and invasion assays. C4-B2-
Infr2shRNA cells expressing inhibited levels of Infr2 and higher levels of Nfr2
showed fewer colonies in soft agar and proliferated faster but did not migrate as
compared to control. C4-B2-Control shRNA cells. Similarly, LNCaP-Nfr2-shRNA cells ex-
pressing inhibited levels of Nfr2 in less metastatic LNCaP cells demonstrated sig-
nificantly higher number of colonies in soft agar, proliferated faster and showed
greater migration and invasion as compared to LNCaP-Control shRNA cells. These
results together suggest that lower Nfr2 levels are associated with higher prolifera-
tion, migration and invasion or metastatic progression. Currently, we are investi-
gating the mechanism of the role of Nfr2 in metastasis progression and plan in vivo ex-
periments to test our hypothesis of the association of lower Nfr2 with metastasis
progression in mice.

**2015 Mechanisms of Oxidative Stress Promoted by 1, 4-Diamino-
2-Butanone in Trypanosoma cruzi and Mamalian Cells.**

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The putrescine analogue 1,4-diamino-2-butanone (DAB) is highly toxic to patho-
genic microorganisms, including various fungi and Trypanosoma cruzi. Similar to
other α-aminocarbonal metabolites such as aminocetone and 5-aminolevulinic acid,
DAB exhibits pro-oxidant properties. DAB reportedly undergoes metal-cat-
alyzed oxidation in aerobic medium yielding H2O2, NH4+ ion, and 4-amin-2-
oxoobutanal, a highly toxic α-oxoaldehyde. Administered to mammalian cell cul-
tures, DAB decreases the cell viability which was shown to be associated with
changes in redox balance. Thus, treatment of KRO cells derived from human colon
carcinoma or cultured LL-MK2 Rhesus epithelial cells with millimolar DAB caused
significant decline in cell viability, which was inhibited by pre-addition of catalase,
aminoguanidine (an α-oxoaldehyde trap), N-acetyl cysteine or reduced glu-
tathione. Now we explore the mechanisms by which DAB exhibits pro-oxidant ef-
facts on trypanomastigotes and on intracellular T. cruzi amastigotes. DAB (0.05-5.0
mM) exposure in trypanomastigotes, the infective stage of T. cruzi, leads to a decline
in parasite viability (IC50 c.a. 0.2 mM DAB: 4 h incubation), changes in morphol-
gy, thiol redox imbalance, and increased TcSOD activity. Medium supplementa-
tion with catalase (2.5 µM) protects trypanomastigotes against DAB toxicity, while
host cell invasion by trypanomastigotes is hampered by DAB. Additionally, intracel-
lar amastigotes are susceptible to DAB toxicity. Furthermore, pre-treatment with
100-500 µM buthionine sulfoxime (BSO) of LLC-MK2 potentiates DAB cyto-
xotoxicity, whereas 5 mM N-acetyl-cysteine (NAC) protects cells from oxidative
stress. These data support the hypothesis that redox imbalance, not only the long reported DAB-promoted inhibition of polyamine metabolism, contributes
to its cytotoxicity in both T. cruzi and mammalian host cells.
Ozone (O₃) exposure in both humans and rodents leads to elevations in alveolar and peripheral inflammation, believed to promote endothelial dysfunction. Although pulmonary inflammation is a well-defined response to inhaled O₃, little is known regarding acute effects of O₃ exposure on the mechanisms responsible for impaired coronary vasodilation. We hypothesized that a single, whole-body O₃ exposure will induce pulmonary inflammation and the transference of pulmonary inflammatory elements to the coronary arteries pre-constricted (~50%) with a thromboxane mimetic (U46619), circulating WBCs and coronary artery endothelial function 24 hrs following O₃ exposure (4h at 1ppm) or filtered air (FA). BALF total protein, cell number, macrophage/neutrophil counts and circulating WBC differentials were assessed. Furthermore, experiments assessing the effects of O₃ on the endothelial-dependent vasodilator, acetylcholine (ACh), in coronary arteries pre-constricted (~50%) with a thromboxane mimetic (U46619) in the presence or absence of luminal administration of the superoxide dismutase mimetic (PEG-SOD). Parallel experiments were performed where rodents were injected (i.p.) with inhibitory antibodies directed against lectin-like oxidized low density lipoprotein receptor-1 (LOX-1). Our results demonstrate significant increases in circulating neutrophils, compared to FA controls. Systemic increases in circulating neutrophils were also evident following O₃ exposure. Moreover, ACH-mediated vasodilation was restored in vessels loaded with PEG-SOD and partially restored in rodents pre-treated with the LOX-1 inhibitor. Our data suggest that that O₃-induced pulmonary inflammation contributes to increased ROS-mediated endothelial dysfunction downstream of LOX-1.

2016 Receptor-Mediated Reactive Oxygen Species Contribute to Impaired Coronary Vasodilation following Acute Ozone Exposure in Rat.

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2017 In Vitro Modification of Plasminogen by Methyglyoxal Disrupts Normal Zymogen Activation.

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Reactive dicarbonyls such as methyglyoxal (MG) are present in blood and react with arginines (R) of target proteins, leading to diabetic micro- and macrovascular complications. Normal MG plasma concentrations are estimated to reach 4.5 μM, with these values tripling as diabetic retinopathy progresses. Dicarbonyls irreversibly modify R residues, resulting in a net loss of positive charge via hydroimidazolone formation. Previous shotgun LC/MS/MS proteomics analysis of human serum reacted with MG (18 hrs at 37°C) indicated that plasminogen (PLG) was readily modified at multiple sites. Subsequent studies in our laboratory determined that modification of R561 of PLG resulted in drastic energy-of-interaction changes between PLG and tissue plasminogen activator (tPA). The model revealed that a MG-adduct formed at R561 decreased the interaction energy by up to 419.4 kcal/mol, 164-fold higher than the minimum energy change necessary for an altered interaction. Modified R561 in PLG is of particular biological interest because of its role in the thrombolytic cascade, R561 being the site of cleavage from PLG to plasmin. In an effort to determine functional consequences of this adduction, MG-modified PLG and its activation by both exogenous and endogenous activators was studied. The activation of MG-modified PLG by streptokinase (STK) was altered in both a time (0-60 min) and concentration (1-500 μM MG) dependent manner. Unmodified PLG (90-100 kDa) was cleaved by STK, with the concurrent formation of angiostatin (ANG; 38 kDa) and plasmin light chain (PLC; 25 kDa). In contrast, such effects were diminished in MG-modified PLG, with reduced accumulation of ANG and PLC, and reduction in the disappearance of PLG. Similar studies are underway with tPA and urokinase. The findings indicate that MG-modification of PLG may disrupt the thrombolytic cascade thereby contributing to vascular complications associated with diabetes. (R24DK083948, ABRC, ES016652, T32ES007091, P30ES00694).


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Cellular oxidative stress is an established causative factor in carcinogenesis. Recent research suggests that elevated oxidative stress in cancer cells represents a phenotypic vulnerability that can be targeted by redox intervention. Multiple myeloma is a redox sensitive hematological malignancy characterized by dysregulation of cell cycle progression and checkpoint control. Guided by a phenotypic drug screen for novel leads that display antiproliferative activity through induction of oxidative stress, we have identified the FDA-approved antimicrobial thiostrpeton (T) as multiple myeloma-directed experimental chemotherapeutic. In RPMI-8226 multiple myeloma cells, major hallmarks of hyperproliferation including hyperphosphorylation of retinoblastoma (pRb; Sc70, Ser807/811) and overexpression of e-cmyc and cyclin D1 were fully suppressed at both mRNA and protein levels at submicromolar concentrations of T. Moreover, T-induced G2/M cell cycle arrest was characterized by inhibitory phosphorylation of cyclin-dependent kinase 1 (CDK1; Tyr15), dephosphorylation of histone H3 (Ser10) and cyclin B1 accumulation as revealed by immunodetection. Flow cytometric analysis indicated rapid induction of cytotoxic oxidative stress followed by destruction of the redox-sensitive oncogene phosphatase cdc25C (cell division cycle 25C), known to abrogate the G2/M checkpoint by activation of dephosphorylation of cyclin B-CDK1 (Tyr15). Strikingly, hydrogen peroxide treatment mimicked T-induced destruction of cdc25C. Vice versa, antioxidant pretreatment (N-acetyl-l-cysteine) prevented cdc25C depletion, inhibitory phosphorylation of CDK1, and enrichment of the G2/M fraction, further substantiating the causative role of ROS in T-induced cell cycle arrest. Taken together, these data suggest feasibility of T-based prooxidant intervention targeting the oncogenic redox-sensitive G2/M regulator cdc25C and its downstream effectors for experimental chemotherapy of multiple myeloma.

2019 Selective Detoxification of Hypothiocyanite by Mammalian Thioredoxin Reductase, the Missing Link in Lung Innate Immunity and Antioxidant Defense.

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Hypothiocyanides metabolize thiocyanate (SCN) in the presence of hydrogen peroxide (H₂O₂) producing the thiol-selective oxidant hypothiocyanite (OSCN). OSCN has been implicated in innate immunity through its ability to inhibit the growth of multiple pathogens. Paradoxically, SCN has been shown to protect cells from promiscuous oxidizing species such as hypochlorite (OCI) by a direct reaction forming OSCN, suggesting its selective detoxification by mammalian cells. Mammalian thioredoxin reductase (TR) is distinguished from lower order TRs by its C-terminal selenocysteine (Sec) redox couple and broader substrate specificity. Recombinant rat TR1, recombinant mouse TR2 and purified rat TR metabolized OSCN with high affinity but do not metabolize OCI. Mutant forms of mouse TR2 lacking Sec had significantly higher Km for OSCN. Recombinant E. coli and P. aeruginosa were inactive. Auranofin, a TR inhibitor, decreased OSCN metabolism in 16HBE lysates. 16HBE cells and P. aeruginosa were exposed to an oxidase-peroxidase coupled system that generated a 100 μM steady state exposure of OSCN or OCI for 2 hours. OCI was highly toxic to both 16HBE and P. aeruginosa, but OSCN was only toxic to P. aeruginosa. Inhibition of mammalian TR with auranofin increased OSCN toxicity in 16HBE cells. These findings implicate mammalian TR as the major detoxification pathway for OSCN and suggest a novel mechanism of innate immunity and antioxidant defense in mammals wherein OSCN formation simultaneously results in pathogen inhibition while preventing damage to host tissue. This study was funded by NIH Grant HL084469 and a Cystic Fibrosis Foundation Research Grant.

2020 GRP78 Contributes to the Selective Cytoprotection Afforded by All-Trans-Retinoic Acid against Renal Injury.


Chemical-induced nephrotoxicity is a major cause of acute kidney injury. Pretreatment of LLC-PK cells with all-trans-retinoic acid (ATRA) affords cytoprotection against chemical nephrotoxics. ATRA pretreatment (25 μM, 24 hrs)
protected against p-aminophenol (PAP, 150 μM, 3 hr), iodoacetamide (25 μM, 2 hr), and H₂O₂ (250 μM, 24 hr)-induced cytotoxicity as assessed by the MTS assay. Moreover, ATRA suppressed tumor necrosis factor-alpha (TNF-α, 30 ng/mL, 2 hr)-induced caspase-3 cleavage by 4-fold, indicative of a reduction in apoptosis. In contrast, pretreatment of cells with ATRA had no effect on cisplatin (25 μM, 24 hr)-induced caspase-3 cleavage. The data reveal that ATRA selectively protects against toxins that induce necrotic and death receptor-mediated (extrinsic) apoptotic cell death but not against mitochondrial-mediated (intrinsic) apoptotic cell death. Although the molecular mechanism(s) underlying the cytoprotective effects of ATRA remain unclear, the endoplasmic reticulum (ER) molecular chaperone Grp78 is a pivotal contributor to the cytoprotective/adaptive response to cell stress. Therefore, we subsequently used Grp78 non-inducible cells to explore the relationship between the ER stress response and ATRA signaling. LLC-PK, cells, in which the induction of grp78 expression was disrupted via stable expression of an antisense grp78 RNA (pKASgrp78), were more sensitive to the cytotoxic effects of ATRA itself induced Grp78 in naïve LLC-PK, and PKNEO cells, it was unable to do so in the pKASgrp78 cells. Thus, Grp78 may contribute to ATRA-mediated cytoprotection and ATRA may provide an effective therapeutic intervention for chemical-induced renal injury or other pathological conditions. (ES006694, ES016578).

**2021 Sulfur Dioxide Modulates Oxidative Stress Responses in IL-10-Deficient Mice with Airway Inflammation.**

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**RATIONALE:** Inhaled sulfur dioxide (SO₂) at low concentrations (1-5 ppm) can produce bronchospasm in asthmatics, linked to increased oxidative stress and airway inflammation (AI), potentially due to a deficiency in IL-10 production. We tested this hypothesis in IL-10-sufficient C57Bl6 (C57), and IL-10 knockout (−/−) mice, with experimentally-induced AI.

**METHODS:** AI was induced by sensitization and challenge with ovalbumin (OVA, O/O), with and without inhaled SO₂ (1 ppm for 6 hr/day over 10 days). Bronchoalveolar lavage fluid (BALF) was analyzed for leukocyte counts; a pro-oxidant/anti-oxidant balance (PAB) assay provided an index of oxidative stress, and interleukin-4 (IL-4) levels in lung homogenates were measured by ELISA, as an indicator of AI. Nitrite levels were measured in BALF by Griess assay, as an index of iNOS activity.

**RESULTS:** Basal levels of nitrite were similar among all study groups (O/O-SO₂: 200 ± 30; O/O: 180 ± 30; C57: 40 ± 10; C57-SO₂: 35 ± 10 μM). However, O/O-SO₂ (15 μM) decreased nitrite levels compared to O/O (P<0.05) and C57-SO₂ (P<0.05). A significant effect of SO₂ was seen in IL-10−/− mice (P<0.05). These results were further confirmed by a reduced malate-, glutamate-, and succinate-driven oxygen consumption by all nine statins. In conclusion, a series of clinically relevant statins affected mitochondrial function and viability of C2C12 myoblasts with different potencies. Further research is needed to investigate the link observed between the mitochondrial dysfunction and the inflammatory and the apoptotic hallmarks of statin-induced myopathies.

**2023 Screening and Characterization of Aldehydes on the Basis of Protein Carbonylation.**

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Proteins are easily susceptible to Post Translational Modifications (PTM) where protein carbonylation is the most common type that plays a pivotal role in etiology and/or progression of lethal diseases like neurodegenerative diseases, cancers, aging, diabetes and sepsis. Since, ROS and secondary products of oxidative stress can induce the protein carbonylation and this type of protein oxidation is regarded as a well-known biomarker of oxidative stress with different susceptibility for different amino acids. In this study, we studied the protein carbonylation caused by different aliphatic (formaldehyde to decanal i.e. aldehydes from carbon 1 to carbon 10) and aromatic (benzaldehyde and its derivatives) aldehydes by applying BSA Protein carbonylation assay. Taking into account the ED 50 values of the aliphatic aldehydes, 2mM and 4mM concentrations of aldehydes were used. Among aliphatic aldehydes, only first two aldehydes (HCHO and CH₃CHO) showed the highest protein carbonylation in a dose dependent manner. On the other hand, aromatic aldehydes showed the following trend: 4-Chlorobenzaldehyde (max at 120 mins) > Cinamaldehyde (max at 120 mins) > Benzaldehyde (max at 60 mins). This result was further confirmed by a reduced protein carbonylation. Protein carbonylation increased with increase in dose except in case of cinamaldehyde and benzaldehyde where high dose decreased the carbonylation. So, our result suggests that aliphatic aldehydes can be characterized as less reactive towards protein carbonylation reaction than aromatic aldehydes. Among aromatic aldehydes, cinamaldehyde and 4-chlorobenzaldehyde can be regarded as fast reacting aldehydes, m-tol and p-tol as slowly re-acting with or without lag phase respectively, ortho substituted aldehydes as causing no protein carbonylation.

**2024 Autophagy Is Protective against CYP2E1-Dependent Arachidonic Acid, Buthionine-Sulfoximine, and Carbon Tetrachloride Cytotoxicity in E47 HepG2 Cells through Preventing Mitochondrial Dysfunction.**

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The goal of this study was to evaluate if autophagy promotes or is protective against CYP2E1 toxicity. E47 HepG2 cells express CYP2E1 and control C34 cells which do not express CYP2E1 were treated with arachidonic acid (AA), or buthionine-sulfoximine (BSO), or carbon tetrachloride (CCl₄), in the presence or absence of 3-methyladenine (3MA) or rapamycin. All three compounds reduced cell viability in E47 cells but not in C34 cells. 3MA further reduced cell viability while rapamycin increased it. These results suggested that autophagy is prevented against AA, BSO and CCl₄ cytotoxicity in E47 cells. These compounds induced oxidative stress in E47 cells as TBARS increased and GSH decreased. 3MA enhanced TBARS and decreased GSH while rapamycin prevented these changes. Flow cytometry suggested that these compounds mainly induced necrosis and 3MA enhanced necrosis but not apoptosis; rapamycin prevented the necrosis. These compounds induced mitochondrial membrane swelling, cytochrome c release, reducing mitochondrial ATP production, 3MA enhanced but rapamycin prevented these changes. Mitochondrial ROS protection was enhanced by rapamycin but decreased by 3MA. E47 cells were transfected with Atg7 siRNA which enhanced AA cytotoxicity, increased GSH while rapamycin prevented these changes. The decline in viability was reversed and protective stress was blocked in the presence of rapamycin. The antioxidant N-acetyl-L-cysteine prevented the AA or Atg 7 siRNA-induced oxidative stress and loss of cell viability.
Abundance of Upstream Consensus Regulatory Element Sequences for Nrf2 in Human Glutathione S-Transferase Genes.

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Nuclear factor erythroid-derived 2-like 2 (NFE2L2, or Nrf2) is involved in antioxidant response to cellular stress. In response to oxidative or electrophilic stimuli, Nrf2 binds its target genes containing an antioxidant response element (ARE), upregulating expression of detoxifying/antioxidant enzymes. Numerous studies have demonstrated that some rodent liver GSTs are highly inducible via activation of the Keap1/Nrf2/ARE antioxidant response pathway by prototypical Nrf2 activators such as BHA, sulforaphane and oltipraz. Thus, these chemicals may have therapeutic value based on their putative ability to activate Nrf2-mediated antioxidant pathways. However, human studies in vitro and in vivo have not generally seen a robust induction of GSTs following administration of Nrf2 activators, suggesting that human liver GSTs are less responsive to ARE-mediated induction. To identify the potential basis for a possible inter-species difference in GST inducibility, we used the web-based oPOSSUM software to identify ARE sequences up to 10,000 base pairs upstream of the transcriptional start sites of antioxidant enzymes putatively regulated by Nrf2 in murine and human genomes. Only three out of 19 human GST genes (GSTM2, GSTO2, and GSTZ1) contained an ARE consensus sequence within 10kb of the transcriptional start site, whereas ARE sequences were present in 11 of 17 murine GSTs analyzed. Notably, none of the common, highly expressed human alpha or mu class GSTs contained consensus Nrf2 elements, whereas 2 of 4 alpha class, and 5 of 7 mu class, mouse GSTs contained one or more Nrf2 responsive sequence elements. Our data suggest that inter-species differences in GST inducibility may result from differences in the presence and location of ARE consensus sequences upstream of murine and human GST genes—a conclusion which may have major implications in the clinical setting.

Pharmacological Inhibition of Thioredoxin Reductase 1 Attenuates Hyperoxic Lung Injury by Augmenting Glutathione Synthesis.

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Inflammation and oxygen toxicity increase free radical production and contribute to the development of acute respiratory distress syndrome (ARDS). We have previously shown increased glutathione (GSH) levels in lung epithelial cells in vitro and attenuated adult murine hyperoxic lung injury in vivo following thioredoxin reductase-1 (TrxR1) inhibition with auranofin and aurothioglucose (ATG), respectively. The present studies tested the hypothesis that ATG treatment increases pulmonary GSH levels, decreases lung injury, and improves survival. Adult male mice were given a single IT dose of 0 or 0.375 mg/kg E. coli LPS. After 12 h, an injection of 0 or 25 mg/kg ATG or a combination of ATG and 800 mg/kg buthionine sulfoximine (BSO) were administered, i.p. Mice were then exposed to room air (RA) or >95% hyperoxia (O2). After 3 d exposure, bronchoalveolar lavage (BAL) and lung tissues were collected. BAL protein concentrations were significantly greater in LPS/O2-exposed mice when compared to PBS/RA controls. In LPS/O2-exposed mice, ATG treatment significantly decreased BAL protein concentrations, increased lung GCLM expression, GSH levels, and GSH/GSSG ratios, and improved survival compared to PBS-treated LPS/O2-exposed mice. BSO treatment dramatically decreased the survival of ATG-treated LPS/O2-exposed mice. In summary, ATG enhances GSH levels, decreases lung injury, and improves survival in a GSH-dependent manner in a murine model of ARDS. If enhancement of pulmonary GSH levels can be accomplished via drug-mediated TrxR1 inhibition, which seems to be without detrimental effects, this approach could constitute a novel strategy to improve outcomes in patients with oxidant-mediated lung injury. This work was supported by the National Institutes of Health (R01 HL105365, TET K08HL093365, and LKR R01AT006880).

Aniline Up-Regulates Cyclins via Induction of Oxidative Stress in Rat Spleen.

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Aniline exposure is associated with toxicity to the spleen leading to splenomegaly, fibrosis and a variety of sarcomas of the spleen. In earlier studies, we have shown that aniline exposure leads to iron overload, oxidative and nitrosative stress and activation of redox-sensitive transcription factors, which could regulate various genes leading to a tumorigenic response in the spleen. However, molecular mechanisms leading to aniline-induced cellular proliferation in the spleen remain largely unknown. This study was undertaken to further assess the role of oxidative and nitrosative stress in the regulation of cell cycle proteins (cyclins) following aniline exposure. Groups of male SD rats were treated with aniline (1 mmol/kg/day by gavage), aniline plus N-acetylcytstein (NAC, an antioxidant, 300 mg/kg/day, i.p.), aniline plus aminoguanidine (AG, an iNOS inhibitor, 200 mg/kg/day, i.p.) or aniline plus zinc protoporphyrin (ZP, a heme oxygenase inhibitor, 50 mg/kg/day, i.p.) for 7 days (controls received drinking water only), and mRNA expression of cyclins A, B, D3 and E were measured in spleen. Aniline treatment resulted in significant increases in the expression of cyclin A (7.9-fold), cyclin B (7.3-fold), cyclin D3 (3.7-fold) and cyclin E (5.4-fold) as compared to the controls. Interestingly, all of the three inhibitors significantly reduced the aniline-induced overexpression of cyclins. Specifically, NAC reduced the expression of cyclins A, B, D3 and E by 61%, 48%, 38%, and 41%, AG reduced cyclins A, B, D3 and E by 60%, 63.0%, 62%, and 51%, and ZP reduced cyclins A, B, D3 and E by 30%, 61%, 62%, and 44%, compared to aniline only treated rats, respectively. Our data suggest that oxidative and nitrosative stress play a role in aniline-induced overexpression of cyclins which could be critical in cell proliferation, and may contribute to aniline-induced tumorigenic response in the spleen. Supported by NIH ES06476.

Understanding the Role of UCP2 in Fatty Acid Beta-Oxidation and Drug-Induced Liver Injury.

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UCP2, originally described as a mitochondrial uncoupling protein, has been reported to serve many other functions including regulation of glucose and lipid metabolism as well as regulation of ROS in cancerous cells. We recently reported that under hepatotoxic conditions such as that induced by acetaminophen, UCP2 expression was found to protect against liver damage in a PARP-dependent fashion. Interestingly, polymorphisms in the coding sequence of the UCP2 gene have been found to be associated with obese and/or diabetic individuals further implicating UCP2 in the fatty acid beta-oxidation pathway. However, the precise mechanism by which UCP2 exerts these salutary effects is unknown. In this study, the metabolic functions of UCP2 and UCP2 (A55V) variant were compared using a gas chromatography coupled with mass spectrometry-based metabolomics approach. COS-7 cells were transfected with myc/flag-tagged UCP2 and UCP2 (A55V). After confirming expression of the constructs via Western blotting (no difference was observed between UCP2 and UCP2 (A55V)), the cells were extracted, metabolites derivatized using methoxamine and MSTFA, and the samples ran on an Agilent 5975C Series GC/MSD. Principal component analysis of the data show distinct separation of the UCP2 transfected cells from the control and UCP2 (A55V) transfected cells. Among others, palmitic acid was found to be dramatically decreased (~2.3 fold) in the UCP2-transfected cells compared to control and UCP2 (A55V) transfected cells. Further, palmitic acid is thought to induce UCP2 expression through PARP activation. This study illustrates the utility of the metabolomics approach for elucidating and/or clarifying the metabolic function of UCP2.

Real-Time Monitoring of Xenobiotic-Induced Intracellular Redox Changes Using Ozone As a Model Oxidant.

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Oxidative injury is often cited as a key feature in the toxic action of many xenobiotics; however, unambiguous indices of xenobiotic-induced oxidative stress have proven elusive. A new generation of sensors capable of reporting intracellular redox...
status has shown much promise. However, their use in toxicological assessments involving strong chemical reagents remains tenuous to O(3), a highly reactive oxidant gas, induces pulmonary function decrements and inflammatory responses in the airways through oxidative mechanisms. An important unresolved toxicological question concerns the effect of O3 exposure on intracellular inflammatory responses in the airways through oxidative mechanisms. An important unsubstantial increase in EGSH in BEAS-2B cells that is indicative of an oxidant-dependent impairment of redox homeostasis. Moreover, this study demonstrates the utility of using redox reporters in making reliable assessments of cells undergoing exposure to xenobiotics with potent oxidizing properties. THIS ABSTRACT OF A PROPOSED PRESENTATION DOES NOT NECESSARILY REFLECT EPA POLICY.

2030 Formation 4-Hydroxynonanoyl, an Electrophilic Lipid Peroxidation End Product, in Rabbit Corneal Organ Cultures Treated with Nitrogen Mustard.

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Sulfur mustard (SM, bis (2-chloroethyl) sulfide) and nitrogen mustard (NM, mechlorethamine) are cytotoxic vesicants that can cause ocular injury. The cornea is particularly sensitive, developing opacity and ulcerations following SM exposure. In these studies we used an air–liquid interface (air lifted) organ culture model in which vesicants are applied directly to the central cornea to assess mustard-induced oxidative stress. Three and six hr post treatment (100 nmol NM in 10 μl PBS), the corneal epithelial layer was found to express proteins containing the lipid peroxidation product 4-hydroxynonanoyl (4-HNE). This was associated with expression of the antioxidant hemeoxygenase-1 (HO-1). To assess molecular mechanisms underlying this response, we analyzed the effects of 4-HNE on human corneal epithelial cells (HCEC) in culture. 4-HNE (0.03 mM) was found to cause a time-dependent induction of HO-1 mRNA and protein; optimal expression of the enzyme was evident after 10 hr. HO-1 is known to be regulated by both MAP kinases and phosphatidylinositol (PI)-3 kinase (PI3K)/Akt signaling. Treatment of corneal cells with 4-HNE upregulated the expression of the antioxidant protein inducible nitric oxide synthase (iNOS) and the level of iNOS mRNA. 4-HNE also increased the expression of the transcription factor HIF-1α and the level of HIF-1α mRNA. 4-HNE upregulated the expression of the antioxidant hemeoxygenase-1 (HO-1). These results suggest that 4-HNE is a potent inducer of HO-1 expression and that HO-1 is a critical factor in the protective response of corneal cells to 4-HNE.

2031 Ionizaid-Induced Cellular Challenge in HL-60 Cells: An In-Depth Proteomic Perspective.

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INTRODUCTION: Ionizaid (INH) is the first line therapy to combat tuberculo- sis. However, its use is associated with potent clinical toxicity, particularly liver damage. However, the rare but important side-effect of INH-induced agranulocyto- sis is an idiopathic drug reaction for which the toxicity mechanism is unclear. Since INH forms various reactive metabolites, we hypothesized that intracellular protein modifications by INH could reveal a possible mechanism of INH-induced agranulocytosis. METHODS: HL-60 (promyelocytic leukemia) cells were used for these studies. Trypan Blue exclusion was conducted for cytotoxicity. Immunoblot assays were conducted by using anti-INH and anti-DMPO antibodies to identify INH-covalently bound proteins and INH-induced protein radicals, respectively. Later, these modifications were confirmed by MALDI-TOF MS. For quantitative proteomics, SILAC (stable isotope labelling by amino acids in cell culture). RESULTS: 5 mM INH had no significant cytotoxicity after 48 hrs. Anti-DMPO (protein radicals) showed a prominent protein radical at 50 kDa, however, multiple INH-covalent protein adducts were found by anti-INH immunoblotting. In SILAC experiments we found changes in 101 proteins (very high confidence limits) of which 38 were upregulated and 63 were downregulated. We observed a significant down regulation of ribosomal proteins, transcriptional factors and splicing factors. But at the same time, factors associated with protein stability were up-regulated. Pro-apoptotic signals were counterbalanced by anti-apoptotic signals, and cell growth factor genes. CONCLUSION: The result showed that INH is acutely non-toxic to HL-60 cells, and is unlikely to possess direct toxicity to neutrophils or bone marrow precursors. INH treatment appeared to induce cellular energy conservation through downregulation of protein synthesis and enhanced protein stability. Further pathways will be elucidated once INH-induced protein radicals and covalent adducts are identified.

2032 Attenuation of Experimental Retinopathy of Prematurity by Vitamin A in the Newborn Rat.

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Supplemental oxygen administration is frequently encountered in the treatment of premature infants suffering from respiratory distress. However, hypoxia contributes to the development of chronic lung disease [bronchopulmonary disease (BPD)] and retinopathy of prematurity (ROP). In this investigation, we tested the hypothesis that exposure of newborn rats to vitamin A and hyperoxia would attenuate retinopathy and abnormal neovascularization compared to those exposed to hyperoxia alone. Newborn Fisher 344 rats were maintained in room air or exposed to hyperoxia (95% O2) for 7 days. Some animals were treated i.p. with vitamin A [2 mg/kg], once daily for the first 5 days of hyperoxic exposure. Animals were sacrificed at selected time points after termination of hyperoxia. Retinal vascular densities of flat mounted retinas were assessed. Protein expression of HIF-1α was determined by Western blotting. Oxidative DNA damage in retinal tissue was determined by 32P-postlabeling. Immediately after 7 days of exposure to hyperoxia alone, we observed constricted retinal vessels, compared to those given vitamin A + hyperoxia. Seven to thirty days after termination of hyperoxia alone, the animals displayed formation of abnormal retinal vessels and capillaries, compared to the vitamin A + hyperoxia group. At the 7 day time point, the HIF-1α protein expression in the hyperoxia + vitamin A group was much higher than the hyperoxia alone group. On the other hand, at later time points, the HIF-1α protein levels were higher in the hyperoxia group. Interestingly, oxidative DNA adducts were significantly decreased in the retinas of animals given vitamin A + hyperoxia. Our study supports the hypothesis that vitamin A protects retina from oxygen-induced abnormal neovascularization, and this is the first report that shows oxidative DNA damage to contribute to experimental ROP.

2033 Systemic Nerve Growth Factor Modulates the Transcription of Amino Acid Transporters and Glutathione (GSH) Synthesis in Mice Striatum.

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Nerve growth factor (NGF) is a member of structurally related proteins, named neurotrophins (NTIs), that regulate neuronal survival, development, function, and plasticity. Moreover, NGF is an important activator of antioxidant mechanisms. These functions of NGF are mediated by the tropomyosin-related kinase receptor A (TrkA). There is evidence that NTIs and their receptors are expressed also in vis- ceral tissues. Physical exercise and stress increase levels of NGF in plasma. Using a murine model we have shown that systemic inhibition of GSH synthesis with L-buthionine-S-R-sulfoximine (BSO) increased brain GSH content and induced the transcription of BSO in liver. Murine striatum cholinergic neurons express TrkA receptors that, when we investigated if i.p. injection of BSO or of sodium arsenite (iAs) modulates the transcription of BSO and trkA as well TrkA phosphorylation in mice striatum. Both agents induced the activation of the NGF/TrkA pathway which correlated with an increased transcription of xCT, LAT1, EAAC1 amino acid transporter genes that provide L-cys / L-cys2 to central nervous system and of GCLm which participates in the de novo synthesis of GSH. The inhibition of TrkA phosphorylation by K252a or anti-NGF neutralizing antibody abrogated the BSO and iAs induced transcription of xCT, LAT1, EAAC1 and GCLm suggesting the participation of this pathway in the in vivo antioxidant response at least in striatum. Furthermore, since anti-NGF neutralizing antibodies would not cross the blood- brain barrier, our results suggest that NGF functions as a systemic redox-sensor in both CNS and peripheral tissues and that the NGF/TrkA pathway plays a critical role in the antioxidant response in the striatum of our murine model. Supported by CONACYT 102287.
Mechanical ventilation with supraphysiological concentrations of oxygen (hyperoxia) is routinely used to treat patients with respiratory distress. However, a significant number of patients on ventilators have enhanced susceptibility to infections and develop ventilator-associated pneumonia (VAP). Pseudomonas aeruginosa (PA) are one of the most common bacteria found in these patients. Previously, we demonstrated that prolonged exposure to hyperoxia can compromise the ability of alveolar macrophages (AM), an essential part of the innate immunity, to phagocytose PA. The objective of this study was to investigate potential molecular mechanisms underlying hyperoxia-compromised innate immunity against bacterial infection in a mouse model of PA pneumonia. Here, we show that exposure to hyperoxia (≥99% O₂) led to a significant elevation in levels of airway HMGB1 and an increased mortality in C57BL/6 mice infected with PA. Treatment of these mice with neutralizing anti-HMGB1 monoclonal antibody (mAb) resulted in a reduction in bacterial counts, injury, and number of neutrophils in the lung and an increase in leukocyte phagocytic activity compared to mice receiving control mAb. This improved phagocytic function was associated with reduced levels of airway HMGB1. The correlation between phagocytic activity and levels of extracellular HMGB1 was found that the transgenic group produce less ROS than the wild type group. Total glutathione (GSH) levels and glutathione peroxidase activity are higher in the transgenic mice compared to the wild type group. We measured the mitochondrial membrane potential (ΔΨm) by tetramethylrhodamine ethyl ester (TMRE) and we found that the transgenic group has less Ca²⁺ uptake than the wild type group. The mitochondrial membrane potential (ΔΨm) of the transgenic mice to wild type mice suggest that ErbB2 alters various oxidative stress pathways. Among them are the products of oxygen metabolism by flavoproteins, heme-proteins, Fe-S-center-containing proteins and other oxidation-reduction centers. Nitric oxide synthase (NOS) reduce redox-cycling compounds such as menadione (MD), forming superoxides, which in the presence of nitric oxide (NO) reacts to form peroxynitrite, both of which cause oxidative injury. Therefore, we examined the mechanism of NO for cell survival during MD-induced oxidative stress under NO inhibition using a NOS inhibitor, L-NAME. Dose viability studies performed with various amounts of MD + L-NAME for 24 hrs allowed us to choose 10 μM MD for further studies and also showed that NOS inhibition did not significantly affect the cell viability at any concentration. Increased nNOS protein expression and increased 3- nitrotyrosine formation, a biomarker for peroxynitrite formation was observed with MD and MD + L-NAME over the controls alone by both Western blotting and Immunocytochemistry, suggesting that the cells compensate for decreased bioavailable NO by increasing nNOS. We examined the major antioxidant involved in cellular protection mechanism at 24 hours. Glutathione peroxidase1 protein levels was decreased with MD-L-NAME treatment which was reversed in the presence of NO donor deta-NONOate with MD+ L-NAME. On the other hand, SOD1 and PRDX6 levels remain unchanged. Interestingly, at 24 hrs, GSH assays and stable isoipe labeling of amino acids with cell culture along with LC-MS analysis also reveal that proteins in glutathione system are co-produced at the inflammatory sites react simultaneously with tyrosine and induce synthesis of peroxynitrite (PN) and HOCl formation (respectively) in vitro. A question that follows naturally but never addressed in detail is what happens when PN and HOCl that are co-produced at the inflammatory sites react simultaneously with tyrosine...
residues in proteins. The significance of these combined oxidations on the issue of biomarker utilization in the study by Whitman and Halliwell (Biochem. Biophys. Res. Commun. 258,168–172,1999) wherein it was shown that the 3-nitro-L-tyrosine was lost to some unknown product(s) following oxidation with HOCl. Another important consequence could be that we need additional biomarkers and their validation. Herein, we report the synthesis and characterization of the oxidation product of HOCl reaction with N-acetyl-3-nitro-L-tyrosine ethyl ester (NANTEE), a model for protein-bound 3-nitro-L-tyrosine. When HOCl was a limiting reagent (HOCl < NANTEE), the major product at pH 7.2 was found to be N-acetyl-5-chloro-3-nitro-L-tyrosine ethyl ester (NACNTEE). Following purification (reversed phase HPLC) and characterization ([H]+ 8.7 Hz, 1H), 3.12 (dd, J = 14.1, 5.8 Hz, 1H), 4.16 (q, J = 7.0 Hz, 2H), 4.63 (dd, J = 8.6, 5.8 Hz, 1H) 4.16 (J = 7.1 Hz, 3H), 1.92 (s, 3H), 2.92 (dd, J = 14.1, 8.7 Hz, 1H), 3.12 (dd, J = 14.1, 5.8 Hz, 1H), 4.16 (q, J = 7.0 Hz, 2H), 4.63 (dd, J = 8.6, 5.8 Hz, 1H), 7.60 (d, J = 2.1 Hz, 1H), 7.86 (d, J = 2.1 Hz, 1H), 8.43 (s, 1H), single crystals of NACNTEE obtained in methanol were used for the determination of crystal structure using KappaCCD (charge-coupled device) diffractometer. It was found that the OH group forms an intramolecular OHO hydrogen bond to the amide O atom, linking the molecules into chains terminating of crystal structure using KappaCCD (charge-coupled device) diffractometer.

N—H...O hydrogen bonds to an amide O atom, linking the molecules into chains.

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Crystal structure was determined by KappaCCD (charge-coupled device) diffractometer. The crystal studied was a non-merohedral twin, with a 0.907 N—H...O hydrogen bonds to an amide O atom, linking the molecules into chains terminating of crystal structure using KappaCCD (charge-coupled device) diffractometer. It was found that the OH group forms an intramolecular OHO hydrogen bond to the amide O atom, linking the molecules into chains terminating of crystal structure using KappaCCD (charge-coupled device) diffractometer.

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The opposing orientation of orientation of toxicology experts and select evidence can skew the accuracy and value of testimony in court. Lawyers and judges are acquiring increasing skills in the technical and scientific aspects of toxicology. Scholarly scientific approaches are required to establish whether techniques and methods asserted to be scientifically sound are valid and support justice. An introduction briefly covers topics helpful to understanding what happens to experts in litigation. This includes the law’s theory of expert evidence, rules of evidence and procedure relating to expert testimony, alternative admisibility rules (Frye v Daubert trilogy), the NRC report and Canadian parallels, a law-science syllogism, the adversary process in theory and practice, ethical principles, clashes of ethics, data (especially quantitative), disclosure of evidence helpful to the other side, and for whom the litigation expert works. The toxicology expert must address relevant science in a fair, representative manner and communicate effectively. The use of Bradford-Hill Criteria (1964) has been promoted to address general causation in toxic tests. While comprehensive, it poses challenges including jargon and communication. Recently, a five question approach was introduced to address specific causation. Explanations and examples of causation will be reviewed. Finally, the 2009 National Academies report recommends that forensic sciences look to academic scientific models for reliable practice and technique validation to support court testimony. Lawyers and courts have become more skeptical of unsupported opinions and increasingly aware of how to combat them. Academic toxicology can offer much to advance the legitimacy of admitted evidence, including in the area of forensics. A discussion will include foibles and solutions.

The present study was undertaken to verify the in vivo effects of thiotaurine (TTAU), aulfane compound reported to exhibit antiradical activity in vitro, against ethanol (EtOH)-induced oxidative stress. For this purpose, TTAU was given to male Sprague-Dawley rats (200-250 g) as a single, 2.4 mmol/kg, intraperitoneal (i.p.) dose 30 min before an oral, 4 g/kg, dose of EtOH (40% v/v). The same experiments were carried out with rats treated with the alcohol dehydrogenase inhibitor 4-methylpyrazole (4MP 75 mg/kg, i.p.) and the aldehyde dehydrogenase inhibitor cyanamide (CYN, 60 mg/kg, i.p.), administered 30 min and 60 min, respectively, before EtOH. Control animals received only physiological saline. All the rats were sacrificed by decapitation at 1 hr after EtOH administration, and their blood and livers were collected for the assay of plasma and hepatic levels of malondialdehyde (MDA) and reduced (GSH) and oxidized (GSSG) glutathione and corresponding activities of the antioxidant enzymes catalase (CAT), glutathione peroxidase (GPx) and superoxide dismutase (SOD). EtOH elevated the plasma

(+34%) and liver (+32%) GSH, lowered the plasma (-65%) and liver (-63%) GSH/GSSG ratio, and lowered the corresponding activities of CAT (-86% and -77%, respectively), GPx (-91% and -70%, respectively) and SOD (-45% and -54%, respectively). Both CYN and 4MP attenuated the alterations caused by EtOH, with the latter compound appearing more potent than the former. In contrast, the GSH/GSSG ratio seen with ethanol was lowered further by CYN but raised by 4MP. TTAU was not only more protective than 4MP against EtOH-induced alterations in the liver and plasma but in its presence the protective actions of CYN and 4MP enhanced the present. The result presents indicate that EtOH metabolism to acetaldehyde and beyond influences oxidative stress by EtOH to a greater extent than EtOH itself, and that such an event can be effectively counteracted by TTAU.

Intra-hepatic cholestasis represents a frequent manifestation of drug-induced liver injury in humans. However, the mechanisms involved are diverse and remain poorly understood. Early hepatic effects of chlorpromazine (CPZ), known for years to induce intra-hepatic cholestasis, have been analyzed using differentiated human HepaRG cells. Bile acids (BA) efflux was found to be inhibited as early as after 30min-CPZ treatment as shown by [H3]-taurocholic acid (TA) intracellular accumulation. This accumulation seemed to be mediated by reactive oxygen species generated by CPZ since it was mostly prevented by N-acetyl cysteine co-treatment. In addition, CPZ-induced ROS generation was accompanied by a disruption of the pericanalicular distribution of F-actin as well as an alteration of the inner mitochondrial membrane potential. Following 24 hour-treatment, CPZ inhibited the expression of the two main canaliculare bile transporters BSEP and MDR3. Moreover, NTCP mRNA and activity as well as CYP8B1 mRNA were inhibited, while MRP4 was overexpressed. These alterations indicate that BA uptake and synthesis were decreased, while BA biliary transport was enhanced. These alterations likely represent hepatoprotective mechanisms against BA toxicity. These data support the conclusion that among other mechanisms, oxidative stress plays a major role as both a primary causal and an aggravating factor in early CPZ-induced intra-hepatic cholestasis in human hepatocytes (This work was supported by the European Community, Contracts LIINTOP-STREP-037499 and Predict-IV-202222).

One of the major efforts in nonclinical regulatory informatics has centered on the development and testing of the electronic data submission standard, also called SEND (Standard for Exchange of Nonclinical Data). This standard was developed by the Clinical Data Interchange Standards Consortium’s (CDISC) SEND Team for nonclinical data collected from animal toxicology studies and allows for the electronic submission of tabulated toxicology data in an electronic format. The initial pilot project began in 2003 and was followed by a second pilot in 2007 focused on CDER-regulated projects. The production of the SEND 3.0 Implementation Guide in 2011 was a major step forward for the electronic submission and exchange of standardized data and enablement of data warehousing efforts that better support scientific review and regulatory science initiatives. However, it is recognized that data standards are simply enablers to support the broader goals of better exploring and exploiting diverse data sets and metadata in order to answer important scientific review and regulatory science questions. The range of needs and challenges in the regulatory pharmacology and toxicology field can be daunting but many of the challenges are shared among key stakeholders (e.g., the US FDA, sponsors, and CROs). These challenges, which range from warehousing to analysis to QA/GLP ramifications, are laying the foundation for a compliant exchange of data and opportunities for collaboration through public/private partnerships. This session will discuss the current computational science initiatives at the US FDA and in industry, existing partnerships, challenges and successes, as well as the transformative effect on the CRO model.
During academic training, postdoctoral and graduate students generally are not provided with opportunities for interacting with toxicologists who are involved in risk assessment and regulatory affairs. The educational training mainly focuses on basic sciences or solving mechanistic problems and thus lacks the practical aspects of risk assessment and regulatory preparation. This concern was discussed at the Education Summit in October 2011, which was organized by the Education Committee of the SOT. Dr. John Dowell’s comment that, “toxicology is what we do, but risk assessment is why we do it,” shows the importance for trainees to become aware of both. Unfortunately, when it is time for the trainee to make the decision on what will be the next step in their careers, they are well prepared on what we do, but fall short on why we do it. The objectives of this session are to provide postdoctoral and graduate students with basic understanding of approaches in risk assessment and regulatory affairs in some of the sectors and to educate them about necessary preparatory steps in this field. In this 80-minute Education-Career Development Session, trainees will become more familiar with the routine job of toxicologists outside of the academic setting. Further, there will be a panel discussion on steps that can be taken during graduate school and postdoctoral training to improve the preparation for a career in risk assessment and regulatory fields. Thus, the participants are expected to gain a basic knowledge of risk assessment and regulatory preparation in the life of a toxicologist and how to pursue this field.

**2045 Hydration with Saline Decreases Toxicity of Calcitriol-Injected Mice in Preclinical Studies.**

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**Purpose:** To study the effectiveness of saline injection in reducing the toxicity profile of calcitriol when coadministered in mice.

**Design:** Meta-analysis of published mice studies with calcitriol.

**Methods:** A comprehensive PubMed and ISI web of knowledge search was performed to identify all published case control articles of calcitriol injection in mice, and relevant articles were selected. Using mortality as an end point to study the toxic effects of calcitriol, the relative risk of mortality in mice given saline injections was evaluated for different calcitriol dosages, as well as mouse and tumor types.

**Results:** Coadministration with 0.25 ml of normal saline solution injected intraperitoneally at a lower mortality rate than calcitriol given alone. The calculated relative risk of mortality was 0.1419 (95% CI 0.0093-2.2133, z-statistic 1.393, p-value = 0.1636) when saline is administered with calcitriol compared to calcitriol alone.

**Conclusions:** Hydration with saline is a common practice in patients receiving calcitriol, which appears to be pertinent in experimental studies of mice receiving calcitriol. It is important to note that results of preclinical studies form the basis for decisions in drug use in patient trials. Decreasing mortality in animal experiments will prove to be a meaningful contribution to the field of research. To avoid some mortality due to toxicity, a common practice for investigators was administering less than the target doses or using saline instead of calcitriol-induced mortality in mice. Combination of high mortality in mice and administration of suboptimal dose in certain situations may impede evaluation of the potential therapeutic effects of calcitriol. The same problem is likely to exist with other drugs as well (e.g., chemotherapeutic agents). This preliminary data indicates that supplemental hydration should be kept in mind in evaluating drug toxicology.

**2046 Identification of Behaviorally Active Doses of Morphine and Evaluation of Its Analgesic Effects in the Rhesus Monkey.**

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Pain affects millions of people worldwide and is commonly treated with opioid analgesics, despite considerable safety/risk management concerns. The purpose of this study was two-fold: First, to establish a therapeutic index for morphine by determining the behaviorally active range in an operant responding procedure and second, to work within this dose range to evaluate the analgesic effects of morphine in the rhesus monkey (Macaca mulatta) in a thermal pain tail withdrawal procedure.

For the operant responding procedure, 2 monkeys were trained to lever-press for food on a fixed-ratio (FR30) schedule. The sessions lasted 120 minutes (8 cycles of 15 minutes) during which the rate of responding was recorded. Acute doses of morphine sulfate (0.3 – 3 mg/kg s.c.) were administered at the beginning of the session. For the tail withdrawal procedure, 4 monkeys were seated in restraint chairs and the lower section of the shaved tail was immersed into water at 40, 50, and 55 C. Sessions began with a control tail withdrawal latency determination at each temperature followed by an acute dose of morphine sulfate (0.3 – 3 mg/kg s.c.). Temperatures were presented to the animals in a randomized order at 15, 30, 60, 90 and 120 minutes post-dose. Morphine markedly decreased the rate of operant responding at 1 mg/kg and eliminated lever-pressing at 3 mg/kg. Analgesic effects of morphine in the tail-withdrawal assay were observed at all doses tested: The onset and duration of action were dose-dependent. The minimum effective dose was found to be 0.3 mg/kg, while 3 mg/kg produced the maximal effect in all 4 animals tested. These results suggest that acute morphine has a narrow therapeutic index in the rhesus monkey. The selected acute doses of morphine therefore provide an appropriate efficacy and safety measures for assessing novel opioid and non-opioid based analgesics with high translational validity and relevance.

**2047 Pharmacodynamic Glycemia Effects from Rapid and Long-Acting Insulins Administered at Mealtime to Alloxan-Induced Diabetic Miniature Yucatan Swine.**

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Pharmacodynamic effects from various insulins with known properties in humans were studied in the Yucatan miniature swine for comparative purposes. Yucatan miniature swine (Sus scrofa, at least 3 months, 20-60 kg) were made diabetic with intravenous alloxan and regulated on insulin. Animals were considered diabetic if they became hyperglycemic (≥150 mg/dL) within 2-5 days following induction. All procedures were on overnight fasted animals (no feed or insulin for 18 hrs). Our diabetic mini-swine average baseline bG of 429 ± 84.5 (SD mg/dL) (N=148 measurements) while non-diabetics average 58.7 ± 8.2 (SD mg/dL) (N=238 measurements). For this study, well-known prototypical marketed insulins (ApidraTM, HumalogTM, Lantus®TM, 0.25 or 0.45 U/kg s.c.) were administered given each time, then blood glucose profiles recorded using handheld glucometer devices (One Touch UltraMax®, Lifescan) over the next 8 hrs (rapid-acting) or 24 hrs (long-acting). Venous blood samples were collected from vascular access ports for bG readings. Blood glucose profile data in the diabetic Yucatan generally compared well to published human glucodynamic data for glycemia effects following insulin. These data suggest the Yucatan diabetic model has similar pharmacodynamic responses to the presentation of exogenous insulins for onset and peak effects but not duration for the rapid-acting insulins. The long-acting insulin (Lantus®TM) peaked in swine.
where no peak (same action throughout day) is normally reported in humans. These differences could be due to either the duration of the lasting, the relative high doses of insulin or the use of a small number of animals in this study.

**PS 2048 Toxicity Mechanisms of Anti-Inflammatory Kinase Inhibitors.**


Improving the safety of drug candidates that enter clinical development requires assays that are more predictive of human outcomes. The BioMAP® platform uses human primary cells to model complex aspects of disease and tissue biology and can be applied to better understand biological mechanisms that underlie drug adverse effects. Here we present the analysis of several anti-inflammatory kinase inhibitors, including inhibitors of p38 MAPK, Jak kinase (tofacitinib) and Syk kinase (fostamatinib) tested in a panel of 12 BioMAP® systems covering a broad range of human biology. Specific effects of these compounds in BioMAP® assays modeling aspects of wound healing and vascular biology appear to correlate with certain side effects of these drugs in patients, including skin rash (p38 MAPK), gastrointestinal perforations (Jak), and hypertension (Syk). These in vitro effects may be useful in screening lead candidates prior to testing in animals or humans.

**PS 2049 Evaluation of PBPK Models for Medical Decision Making upon Acute Chemical Exposure: Dichloromethane (DCM) As an Example.**

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Rationale: Assessing the risks of exposure to hazardous chemicals in emergency situations and decision-making on individual medical treatment is not straightforward. We therefore evaluated the use of physiologically based pharmacokinetic (PBPK) models in this context, taking DCM as an example. DCM is toxic by inhalation and can cause mild to serious health effects.

Methods: In a clinical trial approved by the Hospital Medical Ethical Board, six healthy volunteers were individually exposed to DCM for 1 hour. The volunteers had to apply a paint stripper to a surface without an air mask, in a closed room. Eighteen blood samples were drawn (t = 1-6h after post-dose). Whole blood DCM concentrations and percentages of carboxyhemoglobin (% HbCO) were measured using GC. The DCM air concentration in the room was continuously monitored by a Miran 1B2 infrared spectrophotometer. Experimental whole blood DCM concentrations and % HbCO were subsequently compared to the predictions of a previously published PBPK DCM model.

Results: The DCM external dose (area under the exposure-time curves) ranged between 175 and 390 ppm*h (equal to 608 - 1354 mg*h/m3). DCM blood concentrations reached maximum levels at the end of the exposure (t = 1h) and ranged between 0.25 and 5.1 mg/L. Increase in % HbCO was predicted (maxima reached between 2 and 6 h) and ranged between 0.4 and 2.3 %. The predicted DCM blood concentration ranged between 1.55 and 4.20 mg/L at t = 1h and the predicted % HbCO ranged between 3.1 and 4.1 % at t = 1.5 h. The general form of the concentration-time profiles was well predicted, especially in the elimination phase of DCM from blood.

Conclusion: The model was found to fit experimental DCM blood concentrations reasonably well but should be refined for % HbCO. Such a PBPK model might be helpful in case many individuals are acutely exposed to DCM and for whom HbCO monitoring is not available.

**PS 2050 Specificity Protein (Sp) 1 Transcription Factor Modulates Long Noncoding RNA Expression in Liver Cancer Cells.**

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Hepatocellular carcinoma is one of the most prevalent forms of cancer worldwide and it exhibits highly invasive and metastatic properties. Recent studies have shown that of the small proportion of the genome that is transcribed only about 1.4% encode for protein-coding genes. Of the remaining vast majority of transcripts that do not encode protein, long noncoding RNAs (lncRNAs) have recently gained attention because of their pivotal role in disease. lncRNAs are a class of transcripts longer than 200 nucleotides and have been characterized as having both tumor suppression and oncogenic functions in many types of cancers. Although their mechanisms of action remain largely unknown, many lncRNAs are regulated by transcription factors in a tissue-specific manner. Specificity protein (Sp) transcription factors Sp1, Sp3, and Sp4 are overexpressed in many tumors, and regulate expression of genes required for cancer cell and tumor growth, survival, angiogenesis, and immunomodulation. Sp proteins have also been targeted as the targets of many conventional and alternative chemotherapeutic drugs, and therefore further study of their functions is of great interest. In this study, we examined the role of Sp transcription factors in regulating lncRNAs in liver cancer cells. Using HepG2 and HuH-7 cells as models, we investigated the effects of Sp downregulation on the expression of several lncRNAs as well as cell growth and survival. Downregulation of Sp transcription factors by RNA interference or by drugs that target these proteins identified a set of lncRNAs in liver cancer cells that are modulated by Sp transcription factors. Further studies are underway to examine the specific functions of these lncRNAs in liver cancer growth and metastasis as well as their utility as diagnostic biomarkers.

**PS 2051 The Third Trimester: The Critical Phase of the Deteriorating Effects of Cadmium in Pregnancy.**

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Environmental cadmium (Cd) is rising globally particularly in the developing countries. Reports have indicated that the deleterious effects of Cd may occur at lower levels than hitherto thought. The effects of Cd on female reproduction particularly in pregnancy has received only limited measurement and these reports did not delineate the extent of Cd exposure. In utero adverse effects may be critical. One hundred and sixty six subjects (125 pregnant; 35 non-pregnant), were studied. Pregnant subjects were classified into three trimesters: 1st (35), 2nd (35) & 3rd (55). Cadmium, Cu, Zn, Fe, Se, Serum proteins were determined in all subjects. Third trimester subjects were followed until delivery. Cadmium levels were similar in the 1st & 2nd trimesters but significantly increased in the 3rd trimester compared to controls and 1st & 2nd trimesters. Zine level was significantly decreased in the 3rd trimester compared with the 1st & 2nd trimesters. Importantly, Cd was inversely related to Zn. Thirty-two (58%) subjects delivered normal weight babies, 19 (35%) delivered babies with low birth weight (LBW). Four (7%) delivered babies with high birth weight. Women with LBW babies had significantly higher Cd and low Zn levels as well as low BMI. Cadmium, Zn, Se all correlated inversely with neonatal birth weight (NBW). Cadmium and Se strongly correlated inversely with NBW (r = -0.7, p = 0.000; r = 0.31, p = 0.02) respectively. Zinc also correlated directly with NBW. These data suggest that the third trimester with the lowest Zn level also had the highest Cd level (Cd is a metabolic antagonist of Zn). It appears the critical phase that Cd may elicit its toxic effect in pregnancy is the 3rd trimester and low Zn level may be the driving factor. The third trimester is therefore the phase to target in risk assessment, communication and management.

**PS 2052 Ethanol Is a Significant Cofactor in HAART-Induced Hepatotoxicity.**

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Highly Active Antiretroviral Therapy (HAART) has led to a significant increase in the life expectancy of HIV patients; however, there are significant side effects including lipidostrophy and hepatotoxicity. Alcohol abuse is highly prevalent in HIV infected individuals and hence may be a significant negative cofactor in HAART induced hepatotoxicity. The present study examines the mechanisms underlying HAART and alcohol-induced hepatotoxicity. The effects of HAART drugs (azidothymidine, and Indinavir sulphate) in combination with alcohol were examined both in vitro (H4IIEC3- a rat hepatoma cell line) and in vivo. Individual treatments of H4IIEC3 cells with alcohol and AZT showed a certain level of hepatotoxicity which was significantly increased in combinational treatment of alcohol and AZT. These data indicate that alcohol can induce and further enhance HAART-induced cytotoxicity. Alcohol and HAART drug interactions and hepatotoxicity were also assessed in vivo using an animal model of chronic alcohol feeding. Mice were pair-fed liquid diets (Lieber DeCarli) containing 35% of calories as alcohol (alcohol-fed, AF) or as isocaloric maltose-dextrose (pair-fed, PF). HAART treatment groups received AZT (30mg/kg BW) and IDV (50mg/kg BW) by oral gavage for 5 weeks. Animals exposed to both alcohol and HAART developed increased visceral adiposity compared to pair-fed animals, suggesting disturbances in lipid metabolism in these mice. Lipidostrophy was also evidenced by macro and microvesicular steatosis in the livers; elevated liver triglycerides and free fatty acids. Additionally, animals receiving combinations of alcohol and HAART exhibited increased inflammation and greater hepatic neutrophil infiltration.

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Overall, our data demonstrate that alcohol exacerbates HAART hepatotoxicity, and is a significant cofactor in the development of hepatic steatosis and liver injury.

2053 Acute Exposure to Acrolein, a Ubiquitous Environmental Pollutant, and Anti-HIV HAART Medication Leads to Hepatotoxicity.

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Highly Active Antiretroviral Therapy (HAART) is the current treatment for HIV infection. Although HAART leads to a significant increase in the life expectancy of patients with HIV; the prolonged use of HAART causes hepatotoxicity. This has become a significant clinical problem which leads to discontinuation of therapy in turn causing HIV virus reactivation and the development of AIDS. Additionally, environmental pollutants are known to significantly impact general health as well as therapeutic outcome; however their contribution to HAART therapy-induced liver toxicity is unknown. Acrolein is a common environmental, food and water pollutant and a major component of cigarette smoke. It is also produced endogenously via lipid peroxidation and cellular metabolism. The present study examines the potential impact of acute acrolein exposure on the hepatotoxic effects associated with HIV medication. A well characterized human hepatoma cell line (HepG2) model system was used to investigate the combined cytotoxic effects of acrolein along with HAART drug, azidothymidine (AZT). HepG2 cells were treated with various concentrations of acrolein and HAART drugs either individually or in combination. Our results showed that acute exposure to either acrolein or HAART drugs had minimal to no effect on hepatocyte survival. However acrolein exposure enhanced the AZT-induced apoptotic death in HepG2 cells. Acrolein also sensitized hepatocytes to AZT-induced mitochondrial dysfunction, as shown by mitochondrial membrane depolarization and ATP depletion. Notably, acrolein and AZT responsive epigenetic modifications at the FoxL promoter were observed, leading to a marked enhancement in FoxL gene expression, a known death ligand for hepatocyte and liver damage. Overall, the data suggest that exposure to environmental pollutant acrolein has the potential to exacerbate AZT-induced hepatotoxicity, and increase the severity of drug induced liver injury. This work was supported by NIH grants.

2054 Solubility Enhancement Studies for a Potential Cyanide Antidote.

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Present studies focused on the solubility enhancement for the sulfur donor methyl propyl trisulfide (MPTS) to develop an intramuscular formulation for treating cyanide (CN) intoxication. Various FDA approved co-solvents (ethanol, polyethylene glycols (PEG 200, PEG 300, PEG 400) and propylene glycol (PG)), and surfactants (Cremophor EL, Cremophor RH40, poloxarbate 80, sodium cholate and sodium deoxylcholate) and their combinations were applied to enhance the solubility of the lipophilic MPTS. For solubility determination GC-MS methods were developed. The maximum solubility of MPTS was found at 90% ethanol of over 170 mg/ml. The maximum solubility of over 40 mg/ml was achieved with 20 % Cremophor EL. The combination of the surfactant 20% Cremophor EL and the co-solvent 75% ethanol lead to a synergistic solubilizing effect with the solubility reaching over 400 mg/ml of MPTS.

The in vitro efficacy studies for the MPTS vs. thiosulfate (TS), determined by measuring the thiocyanate formation spectrophotometrically, showed that MPTS is a significantly superior sulfur donor than TS. Similarly, the preliminary in vivo efficacy studies, determined on a therapeutic mice model and expressed as Antidotal Potency Ratios (APR), the ratio of CN LD50 with and without the test antidote(s), showed that MPTS is superior to TS. The combination of MPTS + TS showed a synergistic effect (APR= 3.6).

2055 Chemical Hazards Emergency Medical Management (CHEMM); Chemical Specific Acute Patient Care Guidelines for Prehospital and Emergency Department/Hospital Management.

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Chemical Hazards Emergency Medical Management (CHEMM) is an online and downloadable interactive tool. It is designed to enable first responders, first receivers, other healthcare providers, and planners to plan for, respond to, recover from, and mitigate the effects of mass-casualty incidents involving chemicals. Content has been developed via NLM staff, CHEMM contractors, and Federal government and non-Federal government subject matter experts (SMEs). CHEMM includes chemical-specific, acute patient care guidelines for pre-hospital and emergency department/hospital management for exposures to selected groups of chemicals. The information is divided into sections for response in the Hot Zone, Decontamination Zone, and Support Zone/Treatment Area, and each chemical page includes chemical specific information on substance identification, rescuer protection, triage, pediatric/geriatric/obstetric vulnerabilities, clinical symptoms, antidotes, and more. The first version of CHEMM was released in mid-2011, with recent efforts including the addition of over 50 new chemicals with the types of information noted above.

2056 Manganese Accumulation in the Brain of Asymptomatic Welders and Its Functional Consequences.

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Manganese, a neurotoxicant that is concentrated in welding fumes, may play a role in neurodegenerative processes such as Parkinsonism. In the present study, we examined possible brain biomarkers of manganese exposure in asymptomatic welders using state-of-the-art MRI techniques and correlated imaging findings with functional measures (neuropsychological tests). Sixteen welders and 16 age- and education-matched controls comprised the current sample. For welders, increased welding hours were associated with higher T1 relaxation rates in amygdala, caudate nucleus, hippocampus, putamen, and orbitofrontal white matter in addition to traditionally reported globus pallidus, reflecting increased accumulation of manganese. The higher hours in welding were also associated with greater T1-weighted intensity values in caudate nucleus and hippocampus for welders. Subgroup analysis of welders who were highly exposed to welding compared to controls with no lifetime exposure to welding indicated that welders had higher T1 relaxation rates in globus pallidus and putamen consistent with current literature. Correlation analyses of MRI results and neuropsychological tests revealed that performance on set shifting tasks (stroop and visual-verbal tests) were negatively correlated with T1-weighted intensity values in orbitofrontal white and grey matter for all subjects. In addition, executive function (the D-KEFS tower test) also was correlated negatively with T1-weighted intensity values in amygdala, caudate nucleus, hippocampus, putamen, and globus pallidus. Moreover, welders with high welding exposure showed decreased working memory performance and associated increased T1-weighted intensity values.

These results suggest that brain regions other than the globus pallidus, e.g., the caudate nucleus and/or hippocampus, also may reflect sensitively manganese exposure. In addition, there may be some cognitive decline associated with manganese deposition in brain.

2057 Ethanol-Induced Reductions of Antimicrobial Peptide LL-37 in THP-1 Cells and BAL Fluid of Ethanol Fed Mice.

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The most common infections plaguing the pulmonary system outside of common viral infections such as Rhinovirus and Influenza are Haemophilus influenzae, Klebsiella pneumonia, Burkholderia repaca, Bordetella pertussis, and Mycobacterium tuberculosis. Respiratory infection incidence has been directly correlated to circulating levels of vitamin D. Vitamin D is required for antimicrobial peptide LL-37 production and function in the lung. As a result, antimicrobial LL-37 production is correlated with respiratory function and health. C57Bl6 mice
were exposed to ethyl-alcohol via the Meadows-Cook model for 6 weeks. Mice were also fed Diallyl Disulfide (DADS) at 1 μM/g of feed, and 4000IU of cholecalciferol in their daily diet. THP-1 human acute monocytic cell line were pre-treated with 80nM alcohol for 24 hours, and treated with 1μM of DADS, and 100nM of 1,25 dihydroxy vitamin D for 6, 12, 24 hours. Eighty μM alcohol exposed THP-1 cells displayed a 53% reduction in the cellular supranatant of LL-37. The 6 hour exposure of DADS attenuated the alcohol induced reduction of cellular supranatant LL-37. The 6 hour treatment of 1,25 dihydroxyvitamin D also attenuated the reduction LL-37 in the supernatant of THP-1 cells. In the BALF of the 6 week alcohol fed mice, 1,25 dihydroxyvitamin D was reduced by 70%. In the in-vivo model DADS exposure completely abrogated the alcohol induced reduction of BALF LL-37. This data displays the propensity of alcohol to affect the ability of THP-1 (pre-monocyte cell model) to produce an antimicrobial peptide LL-37. This may be related to the disturbances in 1,25 dihydroxy vitamin D in the pulmonary epithelium as reflected by reductions of BALF of alcohol fed mice. The ability of DADS and vitamin D to attenuate this alcohol induced reduction of LL-37 in the THP-1 cells, and BALF of chronically alcohol fed mice may assist with novel treatment options for human chronic alcohol related respiratory infection severity and rates.

### 2058 Utility of Intranasal Fentanyl Powder Formulation: Pharmacokinetics-Pharmacodynamics Relationship in Rhesus Monkeys.


The recent survey conducted by the American Pain Foundation revealed that breakthrough pain posed the greatest challenge to the quality of life (QOL) of cancer patients. The fentanyl buccal tablet, a self-administration formulation, is used at present to relieve breakthrough pain rapidly. However, a new formulation that possesses much quicker pain relief is required to improve the QOL. SNBL has developed an intranasal fentanyl powder formulation (TRF) with the applied 3Tico System, an intranasal drug delivery system technology. The purpose of this study was to investigate the utility of TRF by comparing the absorption properties and time of onset of action after administration of TRF with those of the commercial buccal tablet. [Methods] Six male rhesus monkeys with body weights 5.4-8.0 kg were used. The plasma fentanyl levels were analyzed by LC/MS/MS. The tail withdrawal latency (TWL) procedure using 50°C water was conducted to evaluate antinociception of fentanyl following administration (8 μg/kg) of fentanyl in 3 forms: TRF (intranasal), intravenously injectable formulation, and buccal tablet. [Results] Tmax and Cmax after administration of TRF were 12.8 min and 2.6 ng/mL, respectively, and were much quicker and higher than the corresponding values for the buccal tablets (50.8 min and 1.1 ng/mL), TRF prolonged the TWL. The effect continued for 25 min from just after nasal administration, similar to that of the intravenously injectable formulation. The effect of the buccal tablet on TWL was noted only at 45 min after buccal administration, suggesting a slow onset of action. [Conclusion] TRF showed a quicker nociceptive effect due to a more rapid nasal absorption of fentanyl in comparison with the buccal tablet. These results indicated that TRF would be a useful formulation for rapidly relieving breakthrough pain in cancer patients with self-administration.

### 2059 Better Prediction of Immunogenicity of Biopharmaceuticals, Is It Possible?

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A major drawback of biologicals is the possible induction of immunogenicity upon clinical use, that may result in a safety issue and/or a reduction of drug efficacy. Anti-drug antibodies (ADA) are determined as a measure of immunogenicity. Current preclinical models have proven lack of predictivity for clinical immunogenicity. Therefore, there is a need for better methods to predict which drugs are likely to induce ADA in clinical trials. Based on historical data of immunogenicity, information on structurally, therapeutically and/or ‘mode-of-action’ similar compounds can be obtained to establish the translational aspects these models. We selected interferon alfa, beta and TNF-inhibitors as model compounds (13 in total) for which the public domain (FDA reviews, BLA, NDA, EMEA, pubmed) was scavenged for immunogenicity related information. The Information consisted of physical chemical properties, formulation aspects (including stability data), preclinical animal toxicity data, clinical data (study specifics and ADA occurrences and effect).

Due to the lack of information on in vivo immunogenicity in literature, we decided to generate these data for 13 selected compounds. We performed autologous ex vivo human T-DC co-cultures and established T cell proliferation and DC maturation as response markers. A logistic regression model has been developed relating physical chemical, animal and in vitro information to immunogenicity, where immunogenicity has been defined as two dimensional: the prevalence in the general population and the potency of the substance (dose-response sensitivity). In a statistical analysis of the model, together with expert information, important predictive factors for immunogenicity were successfully identified.

In conclusion, retrospective analysis of various characteristics and preclinical data from on-market biological can provide us with insight in the mechanism of immunogenicity.

### 2060 Improved Efficacy of Delivery of Antigen Using a Novel Injection Device in the Rabbit.

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Needle-free injector devices show a novel, improved and safer alternative to more classical intramuscular injection immunization protocols. Two routes of administration [intramuscular (IM) versus intradermal (ID) injection; with a gene-gun type needle-free injector device] were compared in rabbits and the immunogenicity response, together with any potential toxicity were evaluated following immunization with a plasmid coding for the Hepatitis B surface protein (HBsAg) when administered up to three times (Days 1 and 29 [and Day 43; intradermal only]). Overall, there were no differences between routes of administration when body temperature, body weight, food consumption, selected csugulation and clinical chemistry parameters, and C-Reactive proteins were compared over a period of 43 days. At necropsy, there were no organ weight changes or any adverse macroscopic observations. Based on the magnitude of antibody response and the number of animals mounting a response, a more robust immune response was observed in animals immunized with the injector delivery method (ID). Only 17% of animals receiving the IM injection showed detectable levels of anti-HBsAg antibodies, compared to 50% of animals receiving the ID injection. Furthermore, anti-HBsAg levels in animals administered ID were generally higher than the IM responder. In conclusion, these results suggest that a needle-free injector device shows a safer alternative to intramuscular injection with an increased efficacy when using in immunization regimens.

### 2061 Development of a Protocol to Determine the T Cell Dependent Antibody Response (TDAR) to KLH in the Mouse—A Comparison between the CD-1 and C57BL/6N Strains.

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Pre-clinical safety assessment of biopharmaceuticals may require testing in a specific strain of mouse, when this is the only species where the test material is pharmacologically active. This can present challenges when evaluating immunotoxicity due to known differences in immune responses that exist between different strains. In this study we developed a robust TDAR protocol in the CD-1 mouse then attempted to transfer a similar protocol for use in the C57BL/6N strain.

Initially CD-1 mice were immunised twice intravenously, 1 week apart, at KLH dose levels of 0.3 or 80 μg/kg/dose. Cyclosporin was also tested at 20 and 100 mg/kg/day in order to establish a positive control. KLH and cyclosporin dose levels of 80 μg/kg and 100 mg/kg respectively, were not tolerated resulting in several premature decedents. A strong IgM and IgG response was observed at the 0.3 μg/kg dose level. Cyclosporin administration at 20 mg/kg/day was sufficient to suppress this response in females only. Cyclosporin administration was tested at 60 mg/kg/day, which proved adequate to suppress the immune response to KLH in both sexes. This protocol was tested in C57BL/6N mice, but failed to produce a robust antibody response. Based on a protocol provided by another lab, C57BL/6N mice were challenged subcutaneously with KLH at 0.1 μg/kg in alum on Day 1 then challenged with KLH only at 0.05mg/kg on Day 7. Control group mice were tested in parallel, without the use of alum. A strong IgM and IgG anti-KLH response was noted in both sexes, most notably in females. In the control group, only the females produced a robust anti-KLH response.

In conclusion, we have established a robust immunisation protocol to measure the TDAR response in CD-1 mice and a suitable positive control to suppress the response. An immunisation protocol in the C57BL/6N mouse was also established.
The T-cell dependent antibody response (TDAR) is a functional assay used in immunopharmacology and immunotoxicology to assess the ability to mount an antibody (IgM and/or IgG) response to immunization. Keyhole limpet hemocyanin (KLH) is extensively used as immunogen of choice both in nonclinical and clinical settings. Native KLH is comprised of high molecular weight (HMW) assemblies (>600-800 kDa) of KLH subunit dimers. It is not known how the different forms (HMW vs. subunit) and manufacturing processes (commercial sources) may impact the nature of anti-KLH immune responses (e.g., magnitude and inter-animal variability). Anti-KLH IgM and IgG responses were studied in female Sprague-Dawley rats immunized on days 1 and 22 with 100 μg of HMW KLH from two different sources or subunit KLHs from three different sources without co-injection with any adjuvant. Dose analysis and biological characterization of KLH formulations were conducted. Anti-KLH IgM and IgG responses were measured on days 1, 8, 15, 22, 29, 36, and 43 using a proprietary indirect electrochemiluminescence immunoassay. The two HMW KLH preparations showed a greater number of sub-visible particles (2-150 μm size range) than the three subunit KLHs. All HMW KLHs and all subunit KLHs were equivalent on SEC (hydrodynamic volume), PAGE (size and charge) and SDS-PAGE (molecular radius). Robust primary and secondary anti-KLH IgG and IgM responses were detected for both sources of HMW KLH. The subunit KLH immunizations resulted in lower IgG and IgM responses, with the exception of Stellar Biotechnologies subunit KLH which produced a robust secondary response. Inter-animal variability for IgM and IgG responses was lower with HMW KLH than with subunit KLHs. In conclusion, different forms and sources of KLH are associated with different magnitudes and inter-animal variabilities in IgM and IgG responses, a critical finding to take into consideration when designing TDAR studies for proper immunotoxicology or immunopharmacology testing.

Assessment of T-cell-dependent antibody responses (TDAR) is implemented in nonclinical safety testing to evaluate test article effect on immune function. Robust ELISA-based assays to analyze hepatitis B surface antigen (HBsAg) and tetanus toxoid (TT)-specific antibodies in serum from cynomolgus monkeys immunized with HBsAg and TT were developed. Assay optimization included evaluation of vaccination doses (Engerix B [HBsAg] at 10 and 20 μg, human pediatric and adult doses; TT Adsorbed at 5 floculation units) and dosing regimen, post-immunization blood collection times, capture reagents and concentrations thereof, initial test serum dilutions, detection antibody dilutions, and incubation times. HBsAg- and TT-specific IgG and IgM endpoint titers (EPT) were analyzed pretest and weekly up to 6 weeks post-immunization to evaluate antibody response kinetics to these antigens. In brief, HBsAg or TT (0.2 or 0.03 μg/well, respectively) is adsorbed overnight to a 26-well microtiter plate following by incubation with an initial test serum dilution of 1:5 or 1:50 and titrated 3x to final dilutions of 1:295245 or 1:2952450 for HBsAg or TT-specific antibody analyses, respectively. Bound HBsAg or TT specific-antibodies are detected using a 1:500 or 1:1000 dilution of alkaline phosphatase-conjugated goat anti-human Ig (total, IgM or IgG). After substrate addition and colorimetric analysis, an EPT is calculated. Following primary immunization, peak HBsAg IgM- and IgG-specific responses were observed 22 and 28 to 35 days post immunization, respectively. HBsAg-specific IgG EPT did not generally increase from primary peak response after a second challenge. Since the study monkeys were immunized with TT prior to purchase, a TT IgM-specific response was not detected. TT-specific IgG responses peaked 15 days post-on-study immunization.
2067 Testing the Use of CD107a and IFN-γ as Markers of Cynomolgus Cytotoxic T Lymphocyte Activation.

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Cytotoxic T lymphocytes (CTLs) are key effector cells, in the immune system, that play an important role in anti-viral host defense and cancer immune-surveillance. Non-human primates, especially cynomolgus macaques, are the closest surrogates for detecting immunosuppressive properties of targeted anti-inflammatory therapeutics in human. While qualified methods exist to measure CTL function in humans and rodents, there are few assays currently available for detection of changes in CTL function in response to drug exposure in cynomolgus macaques. Herein, we present an adaptation of a flow cytometry-based human CTL degranulation assay for characterization of cynomolgus macaque CTL function. This is based on detection of lysosomal-associated membrane protein 1 (CD107a) on the surface of degranulated CTL in conjunction with the secretion of IFN-γ which allows for the identification of fully activated CTL. Briefly, whole blood samples from cynomolgus macaques were stimulated in vitro with staphylococcal enterotoxin B (SEB) for six hours, then stained for the detection of intracellular IFNγ and surface expression of CD3, CD4, CD8 and CD107a prior to analysis on a flow cytometer. Whole blood samples from naive cynomolgus macaques (n = 11) were tested to establish a range of dual positive CD107a+IFNγ+ CTL (0.04-1.58% of CD3+CD4+CD8+ T cells) in response to SEB stimulation. Next, a longitudinal study was completed to determine the intra-animal variation of IFNγ+ and CD107a+ responses in 4 cynomolgus macaques by collecting samples at 2-3 week intervals over a two month period. CTL activation in response to SEB stimulation demonstrated a 94% CV of 31-57% over the course of the study. The direct addition of an immunosuppressive compound, cyclosporine A, to WB suppressed IFN-γ secretion but did not degranulation of CTL in response to SEB in a concentration-dependent manner. The ability to quantify the CTL functional responses in cynomolgus macaques may be a useful tool for determining any immunomodulatory effects of drug candidates in preclinical safety studies on this important lymphocyte subset.

2068 Validation of Assays for Phagocytic Function of Polymorphonuclear Neutrophils and Monocytes in Whole Blood from Cynomolgus Macaque.

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Phagocytosis constitutes an essential arm of host defense against bacterial and fungal infections. The process can be subdivided into phagocyte chemotaxis to sites of inflammation, binding to foreign agents, ingestion of opsonized bacteria, and intracellular killing by oxygen-dependent and oxygen-independent mechanisms. We have assessed several of these functions in whole blood from cynomolgus macaque using commercially available kits designed for human use. Tests included in assay precision, inter-assay precision, processed stability, sample stability, and inter-analyte variability. The percent coefficient of variation was lower for granulocytes than monocytes, typically 1% versus 10% or less in the ingestion assay. Respiratory burst following stimulation with phorbol 12-myristate 13-acetate (PMA), opsonized bacteria, or the chemotactic peptide N-formyl-Met-Leu-Phe (fMLP) was within ranges established for human whole blood. The PMA stimulated was high and consistently > 95% above unstimulated negative control values, the fMLP was low (< 3%), and opsonized bacteria induced a high percentage of stimulated granulocytes (>95%) but lesser stimulated monocytes (<60%). The highest CV values occurred in tests of oxidizing monocytes for inter-assay (19%), processed stability (19%), and sample stability (15%), but were otherwise typically below 5%. Overall these results determine the ability to use these assays for analysis of phagocytic function in cynomolgus macaque under GLP conditions.

2069 Flow Cytometry Immunophenotyping of Lymphocyte Subsets in a Large Cohort of Beagle Dogs.

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Since the dog is a major non-rodent species used in regulatory drug safety studies, there is a growing need to generate historical immunological background data to interpret the results of non-clinical immunotoxicity studies in many species. We have initiated the compilation of data on peripheral blood lymphocyte subsets in a large cohort of male and female naïve Beagle dogs of at least 6 months of age. Venous blood samples of approximately 0.5 ml were collected from fasted animals in the morning. Immunophenotyping analysis of B lymphocytes, total T lymphocytes, T CD4+ and CD8+ lymphocytes with commercial BD Pharmingen fluorescent antibody reagents (BD Biosciences, France) is performed using a Navios 10-color flow cytometer with the Navios v1 software (Beckman Coulter, France). Flow count® fluorospheres are added to each sample prior to flow cytometry analysis for direct determination of lymphocyte subset absolute counts. At writing, validated results are available on 40 males and 50 females. Results obtained in males and females, respectively, are presented as percentage of gated cells and mean (± standard deviation) of absolute count of each lymphocyte subset/μl: B lymphocyte percentage (21.3% & 18.5%) and absolute count (604.4±185.5 & 495.5±171.5); T lymphocyte percentage (60.8% & 64.1%) and absolute count (1719.6±599.4 & 1728.5±34.9); T CD4+ lymphocyte percentage (41.0% & 43.6%) and absolute count (1180.4±399.7 & 1159.5±228.1); T CD8+ lymphocyte percentage (12.7% & 12.2%) and absolute count (360.2±136.6 & 324.6±106.0). Stored samples from additional animals (total of 100 animals from each sex) are being processed, which will very shortly constitute the largest cohort of data on peripheral blood lymphocyte subsets in the naïve Beagle dog.

2070 Development of Molecular Classifiers for Distinguishing between True Sensitizers and False Positives in the Local Lymph Node Assay.

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Recent publications have highlighted certain chemicals which yield false positive responses in the LLNA when compared with guinea pig and human data. Toxicogenomics was applied to provide molecular characterization of the LLNA responses with the goal of developing classifiers for predicting true skin sensitizers. Auricular lymph node gene expression responses were evaluated in female CBA mice exposed to equivalent doses of 9 known chemical sensitizers and 9 known chemical non-sensitizers and 9 false positives per the standard LLNA dosing regimen. Lymph nodes were analyzed for 3HTR incorporation on study day 6 and gene expression responses on study days
4 and 6. Subsequent to data filtering, a comprehensive cross-validation model comparison using 84 different statistical classification methods was performed to identify expression-based classifiers for each chemical class (sensitizer and false positives) and time point. The predictive performance of the models was evaluated using five-fold cross-validation. The gene lists from the top performing model for each chemical class and time point were subsequently confirmed and refined using quantitative RT-PCR. The top performing day 4 classifiers had AUC values of 0.904 and 0.926 for sensitizers and false positives, respectively, while the day 6 classifiers showed higher AUC values of 0.962 and 1.00 for sensitizers and false positives, respectively. The optimal day 6 classifier gene list included 44 and 30 genes for sensitizers and false positives, respectively, with 19 overlapping genes. The 55 total genes displayed functional relevance to sensitization and irritation responses and both classifiers are now being evaluated against a series of new chemicals to independently assess model performance. Overall these data highlight the potential utility of molecular classifiers for distinguishing between sensitizers and false positives.

2071 Development of an Ex Vivo BrdU-Labeling Procedure for the Murine Llna.
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The murine local lymph node assay (LLNA) is widely used to identify chemicals that may cause allergic contact dermatitis. Exposure to a dermal sensitizer results in proliferation of local lymph node T cells, which has traditionally been measured by in vivo incorporation of [3H]thymidine (3HTdr). A more recent non-isotopic variation of the assay utilizes bromodeoxyuridine (BrdU) incorporation in vivo. To eliminate the need for animal injections, we developed an ex vivo labeling procedure. BALB/c mice were dosed topically on the dorsum of both ears for 3 consecutive days with vehicle or one of 3 concentrations of eugenol (EUG), cinnamaldehyde (CA), hexyl cinnaminaldehyde (HCA), nickel sulfate (NiS), iso-propanol (IP) or lactic acid (LA). On day 5, lymph nodes were harvested, and single-cell suspensions were labeled ex vivo with 3HTdR or BrdU. Lymphocyte proliferation was determined by scintillation counting or BrdU ELISA, respectively. Concentration-dependent increases in ear thickness and lymphocyte proliferation were observed for EUG, CA and HCA, but none of the doses tested resulted in excessive skin irritation. The results of both the ex vivo 3HTdr and ex vivo BrdU assays correctly identified EUG, CA and HCA as dermal sensitizers according to the criteria outlined in the Globally Harmonized System for Classification and Labeling of Chemicals. As anticipated, non-sensitizers IP and LA did not induce a positive threshold response in either assay. Furthermore, a positive threshold response was not obtained for the false negative chemical NiS in either assay. The results of both ex vivo assays are in close agreement with those of the in vivo BrdU labeling procedure. We conclude that the ex vivo BrdU labeling method offers predictive capacity comparable to the other previously established LLNA protocols while eliminating animal injections and the use of radiotrace. This abstract does not represent EPA policy.

2072 A Proteomics Approach to Elucidate the Role of Nrf2 in Primary Bone-Marrow-Derived Dendritic Cells of Mice upon Activation by Contact Allergens.
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Contact sensitizers are low molecular weight compounds, which can prompt dendritic cells in the skin to subsequently activate naïve T-cells in draining lymph nodes. Often these compounds are electrophilic and lead to activation of the Keap1/Nrf2 pathway. We aimed to further elucidate the underlying molecular mechanism of skin sensitization and to identify putative biomarkers. We used bone marrow-derived dendritic cells (BMDC) from CD43+ progenitor cells from wild-type or Nrf2-/- mice. Cells were treated for 8 h with different contact allergens: cinnamaldehyde (CA), 1-chloro-2,4-dinitrobenzene (DNB), nickel sulfate (NiSO4) or with SDS as an irritant control. Analysis was done using 2D gels and ESI-MS/MS. Protein spots were considered as differentially expressed if they changed at least 1.5-fold (p < 0.05). While treatment with 250 and 400 μM nickel and 100 μM SDS hardly had any effects, CA and DNBC led to significant changes in protein expression. We found 9 and 27 spots upregulated with 50 and 100 μM CA, respectively. For 5 and 10 μM DNBC 14 and 35 spots were upregulated, respectively. Some of these were detected in Nrf2-/- cells, indicating a Nrf2-dependent regulation. More than 90% of the proteins could be identified. Some of those have been already characterized as Nrf2-dependent such as glutathione S-transferases, glutamate cysteine ligase, or catalase. We also identified other proteins involved in oxidative stress, signal transduction pathways, basic cellular pathways and also heat shock proteins. Some of them were not inducible in Nrf2-/- cells and were not previously described as Nrf2-dependent.

2073 Nrf2-Dependent and -Independent Effects of TBHQ on Early T Cell Activation.
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Nrf2 is a transcription factor that is activated by cellular stress from various sources, including oxidative stress and electrophiles. In response to cell stress, Nrf2 upregulates a battery of cytoprotective genes. In addition to this role, Nrf2 has also been shown to have immunomodulatory effects, including anti-inflammatory effects in innate immune cells and effects on T cell differentiation. The purpose of the present study was to determine the role of Nrf2 in cytokine production and other events associated with early T cell activation. The present studies demonstrate that the Nrf2 activator, tBHq, causes a significant decrease in the early production of IFNγ and IL-2 in anti-CD3/anti-CD28-activated splenocytes from both wild-type and Nrf2-null mice. Interestingly, splenocytes from Nrf2-null mice produced increased levels of IFNγ, but decreased levels of IL-2, as compared to those from wild-type mice. In addition to IL-2 and IFNγ, the effects of tBHq on other genes that are upregulated after T cell activation were investigated. In contrast to IL-2 and IFNγ, tBHq caused only modest effects on the upregulation of CD25 and CD69, which are also induced during early T cell activation. Treatment of activated splenocytes with tBHq also inhibited calcium influx. Collectively, the current studies suggest that tBHq inhibits early production of IFNγ and IL-2 and that this effect is at least partially Nrf2-independent. These studies also suggest that Nrf2 promotes IL-2-, but inhibits IFNγ, at early time points after T cell activation. (This work was funded by NIH grant: E5018885.)

2074 Activation of Nrf2 by TBHQ Inhibits IL2 Production, but Not CD69 Expression, in Human Jurkat T Cells.
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Nuclear factor erythroid 2-related factor 2 (Nrf2) is a transcription factor that is activated by cell stress, such as electrophilic and oxidative stimuli. Once activated, Nrf2 translocates to the nucleus and induces the transcription of its target genes, including HMOX and NQO1. Previously, it has been shown that Nrf2 skews CD4+ T cell differentiation in primary murine T cells, however the role of Nrf2 in human T cells is largely uncharacterized. Therefore, the purpose of the present study was to determine the effects of Nrf2 activation upon early events of CD4+ T cell activation, in human Jurkat T cells. Treatment of Jurkat cells with the Nrf2 activator, tBHq (5 μM), significantly decreased both transcript and protein production of IL-2. Similarly, a modest reduction of CD25 induction by tBHq was observed, but CD69 expression remained unaffected. Nuclear translocation of c-fos and c-jun (transcription factors that regulate IL-2) was not affected by tBHq treatment. In contrast, cells pretreated with 1 μM and 5 μM tBHq showed both a delay of and decrease in Ca2+ influx stimulated by anti-CD3/anti-CD28. Collectively, the current studies suggest a differential effect of tBHq on early events of CD4+ T cell activation, and these effects are at least partially due to inhibition of Ca2+ influx into the cell. (This work is supported by NIH grant: E5018885.)

2075 Inhibition of Early Cytokine Production by the Nrf2 Activator, TBHQ, in Human Primary Blood Mononuclear Cells.
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tBHq (tert-butylhydroquinone) is a common food preservative and a known Nrf2 (nuclear factor erythroid 2-related factor 2) activator. Nrf2 is a ubiquitously expressed transcription factor responsive to cellular stress that regulates many cytoprotec-
Staphylococcal Enterotoxin B (SEB) was used to induce ALI in mice. SEB is a superantigen that activates ~20% of T-cells and massive release of pro-inflammatory cytokines leading to the induction of CD4+Foxp3+ T-cells in the lung, indicating a role for regulatory T-cells in the amelioration of SEB induced inflammation. Additionally, SEB exposure led to the induction of miR-182 that caused a downregulation in Foxo1 expression leading to clonal expansion of T-cells. THC treatment however, decreased miR-182 expression indicating that THC might mediate its effects in part through the regulation of miR-182. Cytokine analysis showed that while SEB exposure led to the increase of IL-2 and MCP-1 in the serum and IFN-γ and IL-6 in the Bronchoalveolar lavage fluid (BALF), THC treatment resulted in a decrease in these cytokines. Together, our data demonstrates that THC can rescue mice from SEB induced ALI and death. (Supported in part by NIH grants P01AT003961, R01AT006888, R01ES019313, R01MH094755, P20 RR026848 and VA Merit Award BX001357)

**2079** Quantitative Phosphoproteomic Analysis of the Dynamic Signaling Network Mediating Proinflammatory Response in the Spleen of Mice under Deoxynivalenol-Induced Ribotoxic Stress.

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The trichothecene mycotoxin deoxynivalenol (DON) is a widely-studied model ribotoxin that targets the innate immune system and has public health significance due to its common contamination of human and animal food. Induction of proinflammatory genes in the spleen by DON is known to involve activation of transcription factors mediated by rapid phosphorylation of mitogen-activated protein kinases (MAPKs). To further understand how phosphorylation of proteins leads to the onset of proinflammatory response, stable isotope dimethyl labeling-based proteomics was applied to quantitatively profile the immediate (< 30 min) phosphoproteome changes in the spleen of mice orally exposed to a toxicologically relevant dose of DON. A total of 90 phosphoproteins indicative of novel phosphorylation events were significantly modulated by DON. In addition to critical branches and

**2078** Identification of microRNA That Affect Multiple Pathways of TCDD-Induced T Cell Dysregulation.

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2, 3, 7, 8-Tetrachlorodibenzo-p-dioxin (TCDD), an environmental contaminant, is well known for inducing severe toxicity including immunosuppression. We examined the mechanisms by which TCDD affects T cell responses. Staphylococcal enterotoxin B (SEB), a superantigen, activates ~20% of T cells via Vβ8 T cell receptor, causes a robust release of pro-inflammatory cytokines. We injected C57BL/6 mice with SEB in footpads and treated i.p with vehicle or TCDD. Our studies showed that TCDD treatment of SEB-activated lymphocyte leads to a decrease in Vβ8+ T cells in comparison to vehicle and causes an increased induction of apoptosis in SEB-activated T cells. TCDD led to induction of Foxp3+ T regulatory cells (Tregs) and suppression of pro-inflammatory cytokines, IFN-γ, TNF-α and IL-6 while increasing the expression of the anti-inflammatory cytokine, IL-10. Showing the role of microRNA (miR) in TCDD-induced immune dysregulation, we performed high throughput miR analysis. Following TCDD administration, 36 miRs were up-regulated while 50 were down-regulated. In silico analysis demonstrated that several pathways were affected including induction of apoptosis, cytochrome P450 expression and Treg differentiation. We validated the role of selected miR following vehicle or TCDD treatment of SEB-activated LN cells. miR-31, complementary to the 3'UTR of the target gene Foxp3, was down regulated in the TCDD treated group which is validated by RT-PCR. miR-351 directed against the target gene CYPIA1, was down-regulated in the TCDD treated group which suggests that TCDD acts as an AhR ligand causing increased CYPIA1 expression as well as Tregs induction. miR-21 targets the pro-apoptotic targets, Fas and Fasl was down-regulated, which results in TCDD-mediated toxicity. Our studies demonstrated that TCDD affects miR expression that acts through several mechanisms to cause immune dysregulation. (Supported by NIH grants P01AT003961, P20 RR026848, R01AT006888, R01ES019313, R01MH094755 and VA Merit Award BX001357)
Penicillium mycotoxins (PM) are natural immunomodulatory contaminants that accumulate in grains, crops, fruits, and fermented products, especially during post harvest periods, due to improper storage and harvesting methods. In this study, we examined the effect of genipin on COX-2 gene expression and analyzed the molecular mechanism of its activity in murine RAW 264.7 macrophages. Furthermore, genipin dose-dependently increased the levels of COX-2 protein and mRNA. These results demonstrate that genipin induced COX-2 expression via NF-κB and AP-1 activation. Moreover, genipin increased the luciferase reporter gene activity in cells transfected with a COX-2 promoter. Transient transfections utilizing COX-2 promoter deletion constructs and COX-2 promoter constructs, in which specific enhancer elements were mutagenized, revealed that the NF-κB, C/EBPβ and AP-1, were predominant contributors to the effects of genipin. Together, these results suggest that genipin induces the expression of COX-2 in response to genipin. Further studies are needed to elucidate the potential role of genipin in the regulation of COX-2 expression in murine macrophages.

BCL-6 and SHP-1: Putative Regulators of TCCD-Mediated Impaired Human B Cell Activation.

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The environmental contaminant 2,3,7,8-Tetrachlorodibenzo-p-dioxin(TCCD) is known to cause suppression of humoral immune responses. Evidence from epidemiological studies performed in dioxin-contaminated areas suggest associations between exposure to dioxin-like compounds and increased incidence of non-Hodgkin's lymphoma (NHL) in human subjects. We have observed that TCCD-treatment of CD40L and cytokine-activated human peripheral blood B cells leads to suppression of B cell activation. Hence, in order to elucidate the molecular mechanisms underlying impaired B cell activation by TCCD, we focused on two candidate genes — SHP-1, a protein tyrosine phosphatase that inhibits signaling in...
activated B cells and BCL-6, a transcriptional repressor of B cell activation and differentiation also mutated in NHL. SHP-1 was identified through a genomic analysis of AHR binding in TCDD-treated mouse B cells. To evaluate the potential involvement of SHP-1 in this process, time-course measurements were performed and SHP-1 mRNA and protein levels were induced at day 3 in presence of TCDD.

In several donors, we also observed a TCDD concentration-dependent increase in protein levels of SHP-1. With respect to BCL-6, we observed decreased downregulation of protein levels compared to control cells. When BCL-6 and SHP-1 levels were measured simultaneously in human B cells, an increase in the double positive (SHP-1hi BCL-6hi) population was seen in the presence of TCDD. This increase in SHP-1 and BCL-6 levels was observed in several TCDD-sensitive human donors and the changes were concentration-dependent. Collectively, these results suggest that the regulators, BCL-6 and SHP-1 may be involved in the TCDD-mediated suppression of human primary B cell activation. (Supported in part by NIH R01 ES002520 and P42 ES049119)

2085 Paradox of Epithelial Early Growth Response 1 in Epithelial Inflammatory Signaling under Ribosomal Insults.

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Regulation of gut epithelial NF-κB expression and activity are crucial for preventing overstimulation of pro-inflammatory response following exposure to commensal bacteria. To determine whether the EGR-1 modulates epithelial NF-κB signaling, we investigated the effects of epithelial EGR-1 on responses to bacterial NF-κB-activating lipopolysaccharide (LPS) in intestinal epithelial cells under ribosomal stress. Although nuclear translocation of NF-κB was observed in the cells exposed to LPS, chemokine expression was slightly affected. In contrast, simultaneous exposure to LPS and ribosomal insults decreased epithelial NF-κB activities, but chemokine expression was super-induced. Similar to our previous study, ribosomal insults-induced EGR-1 mediated induction of pro-inflammatory chemokines in the intestinal epithelial cells. Mechanistically, mucosal ribosomal insult-triggered EGR-1 mediated PPARY induction, which blocked NFκB activation by LPS. Taken together, EGR-1 regulates pro-inflammatory NFκB activation by LPS via EGR-1-induced PPARY although EGR-1 is a positive mediator of chemokine induction by mucosal ribosomal insult in gut epithelial cells. (This study was supported by the National Institutes of Health, grant number K08 AI092457)

2086 TCDD-Induced Modulation of Ig Expression in a Human B Lymphocyte Cell Line.

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2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) is a potent environmental toxin known to inhibit immunoglobulin (Ig) gene expression in various animal studies. We have identified the mouse IgH regulatory region (3′IghRR) as a sensitive transcriptional target of TCDD that may mediate inhibition of Ig expression. Interestingly, the hs1.2 enhancer of the human 3′IghRR is polymorphic and has been associated with a number of autoimmune diseases. In contrast to the inhibitory effects of TCDD on the mouse hs1.2, the human hs1.2 is activated by TCDD. Whether this species difference in hs1.2 modulation translates to the 3′IghRR and Ig expression is unknown. The sensitivity of the antibody response to TCDD-induced suppression in animal models suggests that human B-cells could be a sensitive target of TCDD; however, very few studies have evaluated the effect of TCDD on human B-cell function and Ig expression and none have evaluated the human 3′IghRR. The objective of this study was to characterize the CL-01 human B lymphocyte cell line as a potential human cellular model to determine the relationship between the AhR, 3′IghRR, Ig expression and class switch recombination. Our results support expression of a functional AhR in the CL-01 cells. TCDD treatment also induced transcriptional activity of luciferase reporters regulated by each allele of the polymorphic human hs1.2 enhancer. Additionally, the CL-01 cells can be activated to express Ig by ligands for the Toll-like receptor 9 (TLR9) and this activation appears to be sensitive to TCDD-induced modulation. Future studies will evaluate the role of the hs1.2 polymorphism in 3′IghRR activation, Ig expression and CSR and in the effects of TCDD on these processes. Since TCDD represents a large class of chemicals found in the environment, diet, and pharmaceuticals, understanding chemical-induced modulation of the human 3′IghRR and hs1.2 enhancer may provide a clue to the etiology of autoimmune diseases associated with the hs1.2 polymorphism. (Supported by NIEHS R01ES014676)

2087 Comparison of the Effects of Deoxynivalenol and Tributyltin Oxide to That of Model Compounds Inducing Endoplasmic Reticulum Stress, Ribotoxic Stress and T Cell Activation.

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Two common immunotoxins, the mycotoxin deoxynivalenol (DON) and the organotin compound tributyltin oxide (TBT0), were previously studied for their effects on the human Jurkat transcriptome. DON induces ribotoxic stress and both DON and TBT0 induces ER stress, oxidative stress, calcium mediated signaling, NEAT and NFκB pathways, T cell activation and apoptosis. The present study aimed to confirm this finding by comparing the effects of DON and TBT0 on mRNA expression in Jurkat cells to that of positive controls inducing ribotoxic stress (anisomycin), ER stress (thapsigargin) and T cell activation (PHA and ionomycin).

Jurkat cells were exposed for 6 hours to subcytotoxic concentrations of DON, TBT0, anisomycin, thapsigargin, ionomycin and PHA. RNA was isolated and hybridised on Affymetrix U133 Plus 2.0 Arrays. Effects on mRNA expression were analysed on the level of individual genes or on the level of pathways. The gene expression profiles of anisomycin and DON were almost identical confirming that DON and anisomycin both induce ribotoxic stress and act via the same mechanism. Anisomycin and DON upregulated the processes of ribosomal function, RNA biosynthesis, T cell activation and apoptosis. Genes were similarly affected by thapsigargin and TBT0 confirming that both compounds induce ER stress. Both TBT0 and thapsigargin upregulated genes involved in RNA biosynthesis, ER stress, T cell activation and oxidative stress, and downregulated ribosomal function. Another group of genes were upregulated by TBT0 and not affected by the other compounds and are involved in DNA packaging and nucleosome assembly. As expected, ionomycin induced genes involved in T cell activation. In contrast, PHA did not affect any pathway. In conclusion we showed that DON induces ribotoxic stress and TBT0 induces an endoplasmic reticulum stress response.

2088 Bidirectional Impact of Atrazine-Induced Elevations in Progesterone (P4) on the LH Surge in the Ovariectomized (OVX), Estradiol (E2)-Primed Rat.

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Multiple daily exposures to the herbicide atrazine (ATRZ) have been reported to suppress the luteinizing hormone surge (LHS) in female rats. Exposure has also been found to elevate P4 concentrations, and an increase in P4 is known to have a different directional effect on LH depending on its temporal association to the surge. Consequently, the present study focused on the effects of ATRZ dose and exposure duration on the LHS in OVX, E2-primed Long-Evans rats. ATRZ was administered by gavage (300h), and serial tail blood samples were taken at 1400, 1600, 1800 & 2000h following 1, 2 & 4 days of exposure. An initial study with 0 & 100 mg/kg (previously demonstrated to suppress the LHS after multiple treatments) significantly enhanced the afternoon LH surge peak & area under the curve (AUC) in response to a 1-day exposure, an effect consistent with kisspeptin genetic expression in the brain anterolventriculair periventricular region. In contrast, 4 daily treatments caused a significant suppression in both measures, whereas no effects were present following 2 days of dosing. After a 1-day treatment, prompt & marked elevations in serum P4 concentrations, attributable to adrenal secretion, were present at both 30 minutes & 1 hour, but by the 4th day a marked decline was present at 1 hour. A dose-response assessment was subsequently conducted with 0, 10, 30 & 100 mg/kg ATRZ for 1 & 4 days. At 1 day, 100 mg/kg caused similar elevations in circulating P4, the LH peak & AUC, whereas 4 daily exposures resulted in a shift to a reduced AUC. No effects on the surge were observed with 10 or 30 mg/kg at 1 or 4 days. This influence on the LHS indicates that the effectiveness of ATRZ in inducing P4-associated bidirectional shifts in LH depends on both the dosage administered and importantly on the temporal association of exposure to the appearance of the surge. (This abstract does not represent EPA policy.)

2089 Chlorophyllaziridine: An Ovotoxic Metabolite of Cyclophosphamide?

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Phosphoramide mustard (PM) has been implicated as the ultimate ovotoxic metabolite of the chemotherapeutic agent cyclophosphamide, however, studies suggest that in extra-ovarian tissues PM can spontaneously metabolize to a volatile...
cytotoxic compound, chloroethylaziridine (CEZ). To our knowledge, CEZ toxicity has not been characterized directly in the ovary. Postnatal day 4 (PND4) Fisher 344 (F344) rat ovaries were cultured in media containing vehicle control (1% DMSO; CT) in the absence or presence of PM (60 μM) or CEZ. CEZ-treated ovaries were those that were present in the same incubator as PM-treated (60 μM) ovaries, thus receiving exposure to the volatile metabolite. Additionally, the requirement of ovarian tissue for CEZ formation was evaluated by adding PM (60 nM) to wells that did not contain an ovary to determine any ovotoxic impact on ovaries that were in control media in the same incubator. Following 6 days of culture, follicle types were classified and counted in all treatments. Relative to control, PM and CEZ caused primordial follicle loss (< 0.05), with PM being more ovotoxic (P < 0.05) than CEZ (CT – 42.67 ± 28.88; PM – 86.33 ± 9.9; CEZ – 260 ± 30.0). Small primary follicle depletion only occurred following PM exposure (P < 0.05), not CEZ, relative to CT (CT – 103.3 ± 16.9; PM – 3.67 ± 1.45; CEZ – 83.67 ± 7.8). In the absence of ovarian tissue, CEZ spontaneously arose from PM, and depleted (P < 0.05) both primordial (CT – 377.5 ± 53.24; CEZ – 82.79 ± 29.93) and small primary follicles (CT – 85.0 ± 21.91; CEZ – 24.5 ± 6.6) relative to control. Thus, ovarian tissue is not a requirement for CEZ generation from PM. This study suggests that CEZ is a novel ovotoxicant that warrants further characterization in the ovary to understand its contributions to the detrimental effects of chemotherapy on female fertility. The volatility and toxicity of CEZ is particularly concerning for chemotherapy patients and their families as well as the medical professionals caring for these patients (Supported by ES016818).

2090 Role of Connexin Proteins during the Ovarian Response to 7,12-Dimethylbenz[a]Anthracene Exposure in Rat Ovaries.

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7,12-dimethylbenz[a]anthracene (DMBA) is an ovotoxic polycyclic aromatic hydrocarbon liberated through burning of organic matter. Connexin (CX) 43 and 37 are gap junction proteins providing granulosa-granulosa and granulosa-oocyte communication, respectively, and are essential for follicle survival. CX43 can inhibit chemical-induced apoptosis and low CX43 expression correlates with increased granulosa cell apoptosis. An acute single DMBA exposure to cultured postnatal day 4 (PND4) F344 rat ovaries caused loss of both large primary and secondary follicles at a concentration of 12.5 nM DMBA, while a higher concentration, 75 nM, resulted in only large primary follicle loss 8 days after exposure. This study investigated CX protein involvement during the ovarian response to DMBA exposure. PND4 F344 rat ovaries were cultured for 4 days followed by single exposures of vehicle control (1% DMSO) or DMBA (12.5 nM or 75 nM) and the culture was maintained for 4 or 8 days for Cx43 and Cx37 mRNA or protein quantification (n = 3; 6-10 ovaries/pool and localization (n = 3). After 4 days, Cx37 mRNA was increased (P < 0.05) by 12.5 nM (3.5-fold) and 75 nM (1.5-fold) DMBA exposures, while, conversely, Cx43 was decreased (12.5 nM - 0.94-fold; 75 nM - 0.7-fold; P < 0.05). After 8 days, Cx37 and Cx43 mRNA decreased (12.5 nM - 0.99 and 0.95-fold; 75 nM - 0.85 and 0.95-fold; P < 0.05), regardless of DMBA concentration, compared to control. Cx43 protein was located between the granulosa cells of all follicle stages, while Cx37 was present in the oolemma. After 8 days, decreased (P < 0.05) Cx37 (12.5 nM - 13%) and Cx43 (12.5 nM - 8%; 75 nM - 23%) proteins were observed. These data support an initial increase in Cx37 and Cx43 mRNA as part of the ovarian protective response to DMBA exposure, while the decrease after 8 days of DMBA exposure is likely a reflection of the loss of follicular viability induced by DMBA. (Supported by ES016818 to AFK and AAUW fellowship to SK.)

2091 Endocrine Modulation Potential of Steroids for Possible Applications in Contraception.


The objective of this study was to evaluate novel steroids for selective endocrine modulation potential using high-throughput in vitro screening cell-based and receptor binding systems. Steroids were tested for their potential to bind and modulate activity of the androgen receptor (AR), estrogen receptor (ER), progestrone receptor (PR), and glucocorticoid receptor (GR). For cell-based assays, a reporter gene construct of each hormone response element upstream of the luciferase reporter gene promoter was stably or transiently transfected into mammalian cells and luciferin used for light generation. Relative hormone-responsive activity was quantified by measuring light produced. In the cell-based system, the orally active androgens dimethandrolone (DMA, CDB-1321) and 11β-methyl-19-nortestosterone (CDB-4746) showed androgenic, progestational, anti-estrogenic, and anti-glucocorticoid activities. Their esterified prodrugs, DMAU (DMA-17β -undecanoate, CDB-4521) and 11β-methyl-19-nortestosterone 17β-dodecylcarbonate (CDB-4754) were less progestational and anti-estrogenic than DMA and CDB-4746. In contrast, 17β-unsaturated (CDB-17β-un) decanoate (CDB-3122) only showed androgenic activities. The AR binding system showed that the prodrugs have almost no receptor binding capacity compared with the parent drugs. Levonorgestrel (CDB-0107), a progestational compound used in emergency contraceptive pill, and its esterified prodrug, levonorgestrel-17β-butanone (LB, CDB-1830) showed androgenic, strong progestational and anti-estrogenic, and weak anti-glucocorticoid activities. In conclusion, we have screened steroids for endocrine modulating activity. For esterified prodrugs, liberation of the parent drug by ester cleavage appears essential for biological activity. This work is supported by NICHD Contract N01-HD-9-0014.

2092 Dioxin-Produced Imprinting of Sexual Immaturity through Fixing the Status of a Reduction in the Hypothalamic Expression of Gonadotropin-Releasing Hormone.

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2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD) causes a number of disorders in reproduction and development. Our previous studies have revealed that single treatment of pregnant Wistar rat with TCDD (1 μg/kg, orally) reduces the pituitary synthesis of gonadotropin in perinatal pups, leading to not only a reduction in gonadal steroidogenesis but also the imprinting of sexual immaturity such as defects in sexual behaviors after growing up. However, the reason why the attenuation of pituitary gonadotropin imprints sexual immaturity remains largely unknown. To address this issue, we performed a DNA microarray analysis to identify target gene altered expression of which is linked to dioxin-induced sexual immaturity, using the hypothalami of 70-days-old male pups. The result showed that TCDD causes alterations in the expression of many genes in the hypothalamus (up: 332, down: 283). Among them, we focused on a reduction in gonadotropin-releasing hormone (GnRH) gene, because this hormone plays a pivotal role in the regulation of sexual behaviors. Further analysis indicated that a reduction in GnRH mRNA emerges from postnatal day 4 which is involved in the critical period for the brain organization difference stimulated by sex-steroids. The cerebral level of GnRH protein in the pups was also reduced by maternal exposure to TCDD, although the hypothalamic content remained unchanged. Intracerebroventricular infusion of GnRH into the TCDD-exposed pups after their growing up restored many defects in sexual behaviors. These results together with our previous findings suggest that maternal exposure to TCDD impairs the maturation of GnRH neurons in the offspring by reducing steroidogenesis at fetal and neonatal stage, and this is the origin for imprinted defects in sexual behaviors at adulthood owing to a permanent reduction in GnRH expression.

2093 Bisphenol A Down-Regulates Cytochrome P45011α1 prior to Inhibiting Steroidogenesis in Mouse Antral Follicles.

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Bisphenol A (BPA) is an endocrine disrupting chemical used in polycarbonate plastic products and the epoxy resin lining of food and beverage cans. Once released from these products, BPA readily enters the body and affects physiological processes such as steroidogenesis. Previous studies in our lab have shown that exposure to BPA decreases sex steroid hormone levels in mouse antral follicles. The current study was designed to expand these findings by testing the hypothesis that BPA first decreases the mRNA expression levels of cytochrome P45011α1 (side chain cleave; ScC), which then leads to decreased sex steroid hormone production. To test this hypothesis, antral follicles were mechanically isolated from adult cycling mice, individually cultured in supplemented α-minimum essential medium, and treated with either vehicle control (DMSO) or various concentrations of BPA (1, 10, or 100μg/mL). Follicles and media were collected at various time-points between 6 h and 96 h. Exposure to BPA10μg/mL and 100μg/mL significantly decreased progestrone levels beginning at 24 h and continuing throughout culture compared to DMSO (24h:DMSO:4.86±2.25; BPA10:1.62±0.82; BPA100:0.95±0.06 ng/mL:5-6p±0.05). Exposure to BPA10μg/mL and 100μg/mL significantly decreased androstenedione, testosterone, and estradiol levels at 72 h and 96 h compared to DMSO. Further, exposure to BPA10μg/mL and 100μg/mL decreased expression of ScC by 18 h and that this reduction in ScC leads to a decrease in progesterone production by 24 h, followed by a decrease in androstenedione, testosterone, and estradiol production at 72 h.
Di(2-ethylhexyl) phthalate (DEHP) is a common plasticizer in consumer, building, and medical products containing polyvinyl chloride. Widespread production and use in everyday items represent a public health concern as humans are exposed to DEHP via ingestion, inhalation, and dermal contact. Large doses of DEHP harm ovarian function; however, the effects of DEHP on the ovary at environmentally relevant doses are unknown. Our group has shown that 30 day exposure to relatively low doses of DEHP accelerates recruitment of primordial follicles to the primary stage of development. Immature follicles at this stage rely on intrinsic ovarian factors and proper regulation of the phosphatidylinositol 3-kinase signaling pathway for primordial follicle survival, quiescence, and activation of folliculogenesis. Since DEHP accelerates primordial follicle activation, we tested the hypothesis that 30 day treatment with DEHP alters the expression of intrinsic ovarian factors, specifically those involved in the PISK signaling pathway. To test this hypothesis, CD-1 mice (post-natal day 30) were orally dosed with tocopherol stripped corn oil (vehicle control) or DEHP (20 μg/kg/day, 200 μg/kg/day, 20 mg/kg/day, and 200 mg/kg/day) daily for 30 days. Whole ovaries and ovarian tissue with antral follicles removed were subjected to gene expression analysis by qPCR (n=4/group). In the whole ovary, DEHP increased the mRNA expression of Kit at the 20 μg/kg dose, and decreased the mRNA expression of Pten and Foxl2 at the 20 mg/kg dose and Kit, Pten, Rps6, and Tcfl at the 200 μg/kg dose (e.g. Tcfl vehicle: 1.05±0.19; 200 μg/kg: 0.60±0.02; p≤0.05). In ovarian tissue with antral follicles removed, DEHP decreased the mRNA expression of Mincl at the 200 μg/kg and 20 mg/kg doses. These data suggest that DEHP alters the expression factors involved in early folliculogenesis, specifically in PISK signaling. Furthermore, the dysregulation of PISK signaling could lead to adverse acceleration of early folliculogenesis. Supported by R01 ES019178.

Humoral Immunity in Infant Cynomolgus Monkeys: Control Background Data.

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Background: Developmental immunotoxicity (DIT) evaluations are performed to determine if the test article has an immunotoxic effect on the developing immune system. T-cell dependent antibody response (TDAR) to keyhole limpet hemocyanin (KLH) is one of the methods to assess DIT in the nonhuman primate. Differences in KLH-specific IgM and IgG responses within animal age and animal origin were evaluated in infant cynomolgus monkeys (Macaca fascicularis; hereafter Cynos).

Methods: KLH was administered by intramuscular injection to infant cynos twice. The initial KLH injection (1st dose) occurred postnatal days (PND) 90, 120, 180, or 270. The second KLH injection (boost, 2nd dose) occurred 2 to 3 months after the 1st dose (PND180, 180, 240, and 330, respectively). The origins of the Cynos were Indonesian (Island, IL) and Cambodian (Mainland, ML). Serum samples were obtained prior to each KLH injection and once weekly for 4 weeks following each KLH injections. KLH-specific serum IgM and IgG levels were measured using ELISA methods.

Results: IgM elevated rapidly after the 1st and 2nd doses. The highest IgM in IL (69.2 and 73.6 μg/mL) at the 1st and 2nd doses, respectively) was 24-27% lower than ML (95.2 and 96.4 μg/mL). IgG elevated gradually after the 1st dose but rapidly after the 2nd dose. The highest IgG after the 1st dose in IL (54.5 μg/mL) was 24% lower than ML (45.2 μg/mL). After the 2nd dose, the highest IgG in IL (1339.7 μg/mL) was 39% higher than ML (941.7 μg/mL). The profiles (patterns) of IgM and IgG were similar between IL and ML. IgM was clearly low on PND120/180 group when compared with PND120/180 while IgG was comparable. There were no notable differences in IgM and IgG between PND120/180 and PND180/240 groups. Both IgM and IgG were highest in PND270/DB330 group at most time points.

Conclusion: There were no remarkable differences in TDAR results within animal origins. It is recommended that TDAR assessment be conducted in 4 month or older infants.
2098 Toxico logical Profile and Tolerability of Pixantrone (Pixuvri) in Newborn Mice Comparative Study with Doxorubicin (DOX).

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Pixuvri at the doses of 15 and 27 mg/kg/day was given intraperitoneally to juvenile mice on Post Natal Days 10, 13, 17, 20, 35, 39 and 42 to assess its toxicological profile. Particularly, heart, bone marrow, liver and kidneys toxicity was evaluated for early and late onset.

DOX at 3 mg/kg/day was given as a comparator. Animals were sacrificed at the end of treatment and after 4- or 8-week of observation (DOX terminated after 4-week). Pixuvri up to 27 mg/kg/day was better tolerated than DOX at a dosage nine times lower (3 mg/kg/day) DOX induced a marked reduction in bodyweight gain and a cumulative mortality higher than 50% in pups, survivors being however sacrificed pre-term due to severe decay of their health conditions. Bone marrow toxicity was comparable between Pixuvri at low dose and DOX, whereas no pronounced effects were observed with Pixuvri 27 mg/kg/day, but recoverable after 4 or 8 weeks off-dose. Pixuvri was measurable in plasma up to 2 and occasionally to 6 h, after administration. After repeated treatments Cmax and AUC increased proportionally with the dose and no accumulation was seen. No significant gender difference was observed. Toxicity to thymus and reproductive organs was observed with both test items while no nephro- or hepatotoxicity was detected. Cardiotoxicity was negligible up to 27 mg/kg/day of Pixuvri in females, and quoted as minimal in high dose males at 4- and 8-week of recovery. The cardiotoxicity of DOX, assessed at the end of treatment and 4-week after, was lower than that of Pixuvri, although a significant reduction in heart weight was observed at the end of treatment sacrifice. Notably, for DOX it was not possible to assess the onset and severity of cardiotoxicity at the last timepoint of 8-week, that is definitely the most indicative of late cardiotoxicity. In conclusion, Pixuvri was better tolerated than DOX in newborn and young animals, all reaching adulthood, suggesting Pixuvri as a possible alternative to DOX for pediatric use.

2099 Murine Uterine Receptivity Markers Are Affected by Particulate Air Pollution in a Dose Response Manner.

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It is of concern that exposure to air pollutants (AP) negatively affects reproductive function. Recently, we observed that AP is associated with increased rates of implantation failures (IF) in mice. The uterus requires a subtle collaboration of a variety of factors including cytokines (LIF), adhesion and antiadhesion (MUC-1) molecules and hormonal induced morphological changes (e.g. decidualization, epithelial changes) to become receptive. The aim of this study was to investigate if increased IF due to chronic exposure to PM2.5 could be related to changes in uterine receptivity. To test this, female mice (n=10/group) were continuously exposed (45 days) to either filtered air (AP) or 2 different daily exposure doses of concentrated ambient particles (600 and 1200 µg/m3) of PM2.5. Estrus cyclicity was evaluated 2 weeks before mating and at 4.5 dpc females were euthanized and the following outcomes were evaluated: ovarian and uterine weight, number of corpora lutea, uterine histopathology and LIF and MUC-1 expression by qPCR and immunohistochemistry. We observed that the effects are dose dependent, being more pronounced in the higher dose. Results have shown that estrous cyclicity is affected by a reduction in the duration of the cycle accompanied by an extended diestrus. Ovarian weights are increased but there was no significant change in the number of CL. The histopathological evaluation of the uterus indicated a decrease in the volume and thickness of the endometrium. In both exposed groups, increased diameter and thickness of the glandular and luminal epithelium were seen. No significant alteration was observed in the expression of MUC-1 but there was significant suppression of LIF during the implantation window. In conclusion, exposure to PM2.5 could have significant negative effects on endometrial receptivity by affecting the fine regulation of proliferation and differentiation (decidualization) of uterine cells mediated by LIF expression.

2100 Functional Assessment of Sexual Maturity in Female Cynomolgus Monkey (Macaca fascicularis).

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Selection of suitable criteria for assessing sexual maturity in the female long-tailed macaque (Macaca fascicularis) has yielded conflicting results. The present retrospective work investigates whether the presence of two consecutive menstrual bleedings hallmark complete sexual maturation. Daily vaginal swabs were collected from 1175 Asian and 660 Mauritian origin animals and the records were used to assess the ovarian cycle pattern and seasonality. Animals were housed socially. The swabs were rated from no to heavy menstruation. Menstrual cycle length was categorized (C1, C2, and C3) depending on the cycle length difference between consecutive cycles to distinguish normal cycling females from females with prolonged cycle length. Data from 12446 cycles, comprising 1-38 cycles/animal were correlated with animal age and body weight. Mean cycle duration was 32.4 days. No seasonal differences were detected. The bleeding length for Mauritanian animals (median 2.9 days) was significantly longer than for the Asian animals (median 2.6 days) and the bleeding severity differed also significantly (Mauritian animals: median 3.0; Asian animals: median 2.5). Among the 1835 females under study, 784 had single prolonged cycles and were in the category C2 or C3 whereby the Asian animals had three times more prolonged cycles. At arrival at our facility the average Mauritanian origin animals were significantly heavier (3.75 ± 0.62 kg) than the Asian females (3.25 ± 0.058 kg) although the Asian animals were significantly older (4.74 ± 1.18 years) compared to Mauritanian animals (4.01 ± 0.58 years). This investigation indicates that the onset of sexual maturity is different depending on female cynomolgus monkey origin. However, cycle lengths of the majority of cynomolgus monkeys appeared rather consistent and the cycle category was predictable. This information has to be taken into consideration in the planning phase of studies under the ICH guideline S6(R1) (ICH, 2011) in which the use of sexually mature animals have become part of toxicity studies.

2101 Air Pollution Accelerate Atherosclerosis Progression on Predisposed Mice.

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Exposure to urban particulate air pollution is linked to cardiovascular diseases involving atherosclerosis. In this study we investigated if gestational (G) and/or post-natal (P) exposure to PM2.5 is associated with aortic plaque formation in susceptible individual. LDLr−/− mice were exposed during pregnancy to either filtered or polluted air (daily exposure dose=600 µg/m3 of PM2.5) using a Harvard Particle Concentrator. After weaning, male pups were subdivided and 4 groups were formed according to the exposure period (G or PN). No hypercholesterolemic diet was given to the animals. After 20 weeks of exposure, we assessed non-invasively the size of atherosclerotic plaque (AP) in the aortic arch by ultrasound biomicroscopy. Then, aortic roots were collected and the expression of genes involved in plaque formation and progression was assessed by qPCR (VCAM, ICAM, PCAM, IL-1β, IL-6, IL-10, INFγ, MCP-1, CD36, MMP-1, MMP-9, TIMP-1 and TIMP-2). Birth weight was reduced in animals exposed to PM2.5 during G period (less 11%). Groups exposed to PM2.5 presented greater AP areas and there was an interaction effect (p<0.001) between G and P exposures and the size the plaque (r=-0.43, p<0.01). Expression of IL-1β and IL-6 (proatherogenic cytokines) were higher in exposed groups and IL-10 expression (antiatherogenic cytokines) was reduced. Expression of adhesion molecules and genes involved in plaque destabilization were reduced (VCAM, MMP2, TIMP1 and 2). In most of the cases differences in gene expression were associated with G exposure to PM2.5 but there were also interaction effects with P exposure. For other genes no differences were observed. Results demonstrate how environmental pollution can negatively influence intrauterine environment, impair fetal development and along with postnatal chronic exposure predispose susceptible individuals to atherosclerotic plaque formation later in life. Imbalance between pro- and anti-inflammatory cytokines might account for the progression of atherosclerosis due to PM2.5 exposure.
2102 Brominated Diphenyl Ether-47 Induces Oxidative Stress-Stimulated Pro-Inflammatory Pathways in Human Extravillous Trophoblasts.

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Preterm birth is associated with significant infant morbidity and mortality. Although the etiology of preterm birth is not fully determined, critical roles of oxidative stress and inflammation are implicated. Polychlorinated diphenyl ethers (PBDEs) are widely used flame retardant compounds. Brominated diphenyl ether (BDE)-47 is one of the most prevalent PBDE congeners found in human breast milk, serum and placenta. Despite the presence of PBDEs in human placenta, the effects of PBDEs on pregnancy are poorly understood. The present study investigated BDE-47-induced oxidative stress and the role of oxidative stress on human extravillous trophoblast cell line, HTR-8/SVneo. HTR-8/SVneo cells were exposed to 5, 10, 15 and 20 μM BDE-47 for 4 h and reactive oxygen species (ROS) generation was measured using the dichlorofluorescein (DCF) assay. Inhibition of ROS formation was measured after pre-treatment for 1 h with defereroxine (DFO), an iron-chelating antioxidant. To determine oxidative stress-mediated activation of inflammatory pathways by BDE-47, HTR-8/SVneo cells were pretreated with 1 mM DFO for 1 h prior to BDE-47 treatment for 24 h, or co-treated with 100 μM (α-tocopherol for 24 h. Cytokine release was analyzed by enzyme-linked immunosorbent assay. Treatment of HTR-8/SVneo cells with 15 and 20 μM BDE-47 increased DCF fluorescence compared with solvent controls, indicating increased ROS formation. When cells were pretreated with DFO, BDE-47-stimulated DCF fluorescence was decreased. Pre- or co-treatment with DFO and (α-tocopherol prevented BDE-47-induced interleukin-6 release. These data indicate that BDE-47-induced cytokine release in HTR-8/SVneo cells depended on ROS formation. Because inflammation occurring at gestational tissues during pregnancy has been associated with preterm birth, further research is needed to ascertain potential relevance of these findings to pregnancy and preterm labor.

2103 One-Generation Reproduction Study of Isobornyl Acetate in Rats, with an Evaluation through Sexual Maturity in the F1 Generation.

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Isobornyl acetate, a widely used fragrance ingredient, was administered to male and female rats (25 rats/sex/dose) at dosages of 0, 30, 100, and 300 mg/kg/day. Administration via gavage of the test material or the vehicle, corn oil, began before the cohabitation period (83 days for males; 14 days for females), through cohabitation (maximum 14 days), until the day before sacrifice (males) or day 23 of presumed gestation (females that do not deliver) or day 22 of lactation. F1 generation rats selected for continued evaluation were sacrificed on day 60 (+/- 3) postpartum. There were no treatment-related deaths at any dosage level tested. Excess salivation was observed in P generation male rats at 300 mg/kg/d and P generation females at 100 and 300 mg/kg/d. Mean body weights, body weight gains and feed consumption values were comparable among the dosage groups. There were 23, 22, 24, and 22 pregnant rats in the four dosage groups; all pregnant rats delivered litters. All natural delivery and litter observations were comparable. There were no treatment-related clinical signs, gross lesions or changes in body weight, body weight gains, feed consumption, or organ weights in the male and female F1 generation rats at any dosage level tested. Sexual maturation was unaffected in the F1 generation. Based on the results of this study, the no-observable-adverse-effect-level (NOAEL) for toxicity of isobornyl acetate is 300 mg/kg/day. The reproductive NOAEL in the P generation rats and the NOAEL for viability and growth of the F1 generation offspring is greater than or equal to 300 mg/kg/day. This dose is 2000 times greater than the conservatively calculated exposure (assuming 100% dermal absorption) to isobornyl acetate from fragrance use.

2104 Di (2-Ethylhexyl) Phthalate Is Not Genotoxic to Oocytes, but May Induce Apoptosis in Two-Cell Zygotes.

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Di (2-ethylhexyl) phthalate (DEHP) is a phthalate found in food packaging, toys, medical equipment, among others. DEHP decreases the fertility capacity of the oocyte by interfering with its progression through meiosis and impairing embryo development. Since DEHP is genotoxic to different cell types, we suggest that the decrease in the fertility capacity of the oocyte is also attributed to the genotoxic effect of DEHP. This study evaluated whether in vivo exposure to DEHP causes DNA damage in oocytes, and if this damage persists in the zygote. Female CD1 mice (n = 6 per group) were exposed orally to DEHP (20, 200 and 2000 μg/kg/d) or vehicle (corn oil tocopherol-stripped) every 24 h for 3 estrous cycles. The minimal dose of DEHP was based on the reference dose established by the Environmental Protection Agency (EPA). 5-fluorouracil (1 μM) was used as positive control. Following treatments, mice on estrus received equine chorionic gonadotropin (eCG; 5 IU) ip, 48 h later mice received hormone human chorionic gonadotropin (hCG; 5 IU) and 16 h post-hCG mice were either euthanized to collect eggs or bred with a fertile proved male mouse to collect zygotes. Oocytes were incubated with hyaluronidase to remove cumulus cells and then with Tyrode solution to remove the zona pellucida. Denuded oocytes were used to evaluate viability by staining with propidium iodide/Hoechst 33342 and DNA damage by comet assay. Viability was expressed as percentage live cells and DNA damage was assessed by the Olive Tail Moment (OTM). No significant differences in viability or OTM were observed among groups. Two-cell zygotes from mice exposed to 2000 mg/kg/d DEHP were used to assess the induction of micronuclei (MN) by staining with Hoechst 33342. DEHP did not induce MN in the zygote, however, intra-cytoplasmic microns with cell death by apoptosis were observed. These data suggest that DEHP is not genotoxic to oocytes, but it could elicit apoptosis in two-cell zygotes and thus contribute to reduced fertility. Conacyt-Mexico CB-167/678.

2105 Inhaled Ambient Particulate Matter Induces Preterm Birth and Low-Birth Weight in a Mouse Model of Pregnancy.

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Recent epidemiologic evidence has linked exposure to elevated levels of particulate matter (PM) in air pollution to preterm birth (PTB), low birth weight (LBW), and small for gestational age babies. However, several questions remain unanswered including mechanisms of action (MOA), gestational windows of vulnerability, and most active PM size and composition. To begin to better understand how PM exposure impacts fetal development, timed-pregnant B6C3F1 mice were exposed to a mixture of PM of different PM sizes (CAPs) for 6hr/day, 7days/wk. The PM/m3. Based on EPA standards for PM 2.5 of 35 μg/m3, the exposure for 6 hr/day at our site in Tuexedo, NY or to filtered air (FA) for 6hr/day, 7days/wk. The average daily CAPs concentration throughout exposure was 163.8 ± 99.6 μg PM/m3. Based on EPA standards for PM2.5 of 35 μg/m3/24 hr, our 24 hr time-weighted average (i.e. 6.8 μg/m3) is relevant for many U.S. urban centers. Exposure to CAPs in this study reduced the duration of gestation by 0.5 days (compared to FA controls), which is comparable to a 1 wk decrease in human pregnancy bringing birth closer to the PT category (i.e. <37 wk). Concomitant with the observed PTB were both an 11.4% decrease in birth weight and 2.5% decrease in crown-to-rump length (CRL) at birth and at postnatal day 1. Neovas born to CAP-exposed dams remained lighter than their filtered-air counterparts through the 21 days of nursing, though daily rates of weight gain were equal to that of FA controls. Exposure to airborne PM had no effect on conception rate, maternal weight gain during pregnancy, offspring genotoxicity, or sex ratio compared to FA control. These studies are the first to provide biological plausibility for the epidemiologic evidence demonstrating an association between PM exposure and PTB/LBW. The same recently-developed mouse model will be used in future studies to determine MOA and PM sensitive gestational "windows" of vulnerability. Supported by March of Dimes and NIHES NYU Center ES002060.

2106 In Utero Exposure to Bisphenol A Increases Germ Cell Apoptosis in the Neonatal Mouse Ovary.

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Bisphenol A (BPA) is a known endocrine disruptor that is widely used as a synthetic plasticizer to harden polycarbonate plastics and epoxy resin. However, BPA can leach from plastic products into the food and water of consumers, leading to human exposure to BPA on a daily basis. The objective of this study was to examine the effects of prenatal BPA exposure on early ovarian development in mice. Pregnant dams (6 per treatment group) were orally dosed with tocopherol stripped corn oil (vehicle control) or BPA (0.5 and 50 μg/kg/day) daily starting on gestation day 11 and continuing throughout pregnancy. After birth, neonatal ovaries were collected and subjected to histological evaluations and gene expression analyses. Our results show that the selected doses of BPA affected ovarian development differently. Specifically, compared to vehicle controls, BPA 0.5 μg/kg/day treatment increased the number of healthy germ cells present in germ cell nests and increased SOT 2013 Annual Meeting 449
the number of primordial follicles, suggesting that BPA at this dose prevents apoptosis of germ cells, which is critical for the formation of the primordial pool. In contrast, BPA 50 μg/kg/day significantly increased the number of apoptotic germ cells, indicating BPA at this dose might accelerate germ cell apoptosis. To explore why the selected doses of BPA affect early ovarian development differently, we compared the expression of various apoptotic factors in the treatment groups. BPA 0.5 μg/kg/day decreased the expression of the pro-apoptotic factors Bak1 and Bax, and the ratio of Bcl2/Bax, but decreased the expression of the pro-apoptotic factor Bak2. In contrast, BPA 50 μg/kg/day significantly increased the expression of the pro-apoptotic factor Bak1. Collectively, these data suggest that in utero exposure to BPA affects early ovarian development, but that selected doses of BPA affect ovarian development differently. Supported by: NIH T32ES007326 (WW), NIH P20ES018193 (JAF), NIH K99ES021467 (ZRC).

207 Enhanced Pre-Postnatal Toxicity Study of AMG 162 Administered by SC Injection to Pregnant Cynomolgus Monkeys with up to 6-Months Postnatal Evaluation.


AMG 162 is a fully human IgG2 monoclonal antibody that inhibits bone resorption by targeting RANKL, an essential mediator of osteoclast formation, function, and survival. Monthly SC injection of pregnant cynomolgus monkeys with 0 or 50 mg/kg AMG 162 from ~GD20 until parturition resulted in test article-related effects on body weight gain and their offspring. In infants, there were reductions in biomarkers of bone turnover, increased stillbirths, one instance of dystocia, and one stillbirth following maternal signs of hypocalcemia. Lactation, mammary gland histomorphology, and fetal growth were comparable in controls and the AMG 162 group. In infants exposed in utero, there was increased postnatal mortality, decreased body weight gain, decreased growth/development, and decreased biomarkers of bone turnover from birth to 10 weeks of age. AMG 162-related effects in infants were present in bones (osteoclast hypoplasia, nonproliferative hyperostosis, physeal hypertrophy, and decreased marrow space), normally erupted teeth (dysplasia and malalignment), lymph nodes (most absent), and in multiple tissues (extramedullary hematopoiesis). Signs of infection in multiple tissues were detected in 3 infants that underwent unscheduled necropsy. In necropsies necropsied at 6 months of age, there was full recovery from all bone-related changes observed earlier postpartum with some AMG 162-related effects persisting (absent/reduced size of lymph nodes, extramedullary hematopoiesis, dental dysplasia). One AMG 162-exposed infant, in which there had been recovery from pharmacological effects, had minimal to moderate mineralization of multiple tissues. In general, the effects observed in mothers and infants were consistent with the pharmacological action of AMG 162.

208 Impact of Feed Choice When Performing Generational Reproduction Studies.

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The selection of an appropriate lab diet is extremely important for reproduction (repro) studies. In our lab, Purina Mills (PM) RODENT Diet 5002 has been the standard feed for all rodent repro studies for over 22 years. Our lab has not veered from this diet as a strong historical control exists with its utilization. A new diet, Harlan Teklad 80 (HT) Global 21% Protein Rodent Meal 2016CM, was recently introduced for use in an extended one-generation study. This feed was recommended by the supplier based on its use in other labs for repro studies and it meets the qualifications stated in guidelines of the Endocrine Disruptor Screening Program. Per guidelines, the Genistein-equivalent content of genistein plus daidzein (glycone forms) of each batch of feed must be ≤ 300 μg/g. - 300 – 550 μg/g in the standard diet used in the past studies. During the delivery phase of the study, an increased incidence of pup mortality was noted in all groups including the control group. An increased incidence of pups with no milk in their stomach was also noted in all groups. Upon further evaluation, it was discovered that the feed was occluding the nostrils of many of the pups, thereby interfering with the suckling process. The general physical appearance of the HT feed was very similar to the PM, so feed-related differences due to physical form were not anticipated. However, after discovering the cause of mortalities relating to the feed, it was then noted that the texture of this feed was much finer (similar to talcum powder). Therefore, while a specific feed may be suitable for many repro & developmental study types, the feed may not be suitable for those study types where offgassing are present. While the HT may be an excellent feed source for many repro & developmental study types, we did not consider this feed type to be suitable for our one-generation study.

Although, our findings are based on one experience in our lab, our findings demonstrate that a cautious approach should be taken when making changes in study design as any changes can be crucial to study outcome and interpretation of results.

2091 In Vitro Exposure to Di-n-Butyl Phthalate (DBP) Decreases the Expression of Cell Proliferation Transcripts in Cultured Mouse Ovarian Antral Follicles.


Di-n-butyl phthalate (DBP) is commonly found in consumer products such as plastics, cosmetics, insecticides and oral medications. We have shown previously that DBP (100 μg/mL) inhibited mouse antral follicle growth in vitro. In the present study, the effects of DBP (1000 and 10000 μM) on mouse antral follicle cell proliferation transcripts were determined. Follicles were cultured for 28 days. Males and females were combined for the second half of the exposure (days 14-28) in order to assess whether DBP caused a significant change in steroid production and reproductive physiology. At the end of the exposure, all follicle were treated in vitro. Furthermore, we have shown that DBP may do so by altering the expression of important cell cycle regulators and causing an imbalance in the expression of key members of the Bcl2 family, markers for the intrinsic apoptotic pathway. However, no studies have evaluated the effects of DBP on other cell proliferation markers or the extrinsic apoptotic pathway. The purpose of this study was to further investigate the mechanisms by which DBP inhibits follicle growth and causes antral follicle death. We hypothesized that DBP decreases the expression of cell proliferation genes such as Mek67, Pen, and Igf1, and also increases the expression of Fas, a marker for the extrinsic apoptotic pathway. To test our hypotheses, antral follicles were isolated from adult CD-1 mice (32-37 days old) and individually exposed (n= 8-12/culture) to DBP (1-1000 μg/mL) or vehicle (dimethylsulfoxide, DMSO) for 24 h. Following culture, follicles were subjected to qPCR analysis for the expression of Mek67, Pen, Igf1, and Fas. DBP treatment (100 and 1000 μg/mL) decreased the expression of Mek67 and Igf1 compared to the control follicles (p<0.05). DBP treatment did not alter the levels of Pen transcript when compared to control follicles, while Fas mRNA was undetectable in all treatment groups. Decreased expression of the cell proliferation genes Mek67 and Igf1 further supports that DBP-induced inhibition of antral follicle growth involves a defect in follicular cell proliferation. Also, undetectable Fas suggests that the extrinsic apoptotic pathway may not be involved under these conditions. Supported by NIH grants K99ES021467 (ZRC), R01ES019178 (JAF), T32ES007326 (WW) and the Billie Field Fellowship in Reproductive Biology (WW).

2101 Evaluating Benzo[a]pyrene Effects on Steroidogenesis and Reproduction.

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Benzo[a]pyrene (BaP) is a polycyclic aromatic hydrocarbon (PAH) that has been implicated in modulating aromatase enzyme function. This effect has potential to interrupt normal reproductive function by causing imbalances in homeostatic androgen and estrogen levels. The aim of this study was to use a fish model, Fundulus heteroclitus, in order to assess whether BaP caused a significant change in steroid concentrations that could negatively alter additional reproductive biomarkers. Adult fish were exposed to waterborne BaP concentrations of (0, 1 or 10 μg/L) for 28 days. Males and females were combined for the second half of the exposure (days 14-28) in order to quantitate egg production and fertilization rates. BaP exposure did significantly reduce male gonad somatic index (GSI) and egg fertilization at 10 μg/L. Testosterone concentrations in males were significantly reduced at the high BaP dose averaging only 355 pg/mL plasma versus 2510 pg/mL in the controls. Also, estradiol concentrations in the females were significantly reduced at 1 and 10 μg/L BaP averaging 4070 and 3350 pg/mL plasma respectively compared to 7540 pg/mL plasma in the controls. Sperrn concentrations, egg production, male liver somatic index (LSI) and female GSI and LSI were not altered. BaP exposure at these environmentally relevant concentrations caused negative alterations to both molecular and phenotypic biomarkers associated with reproduction. Our next goal is to assess if these parental effects will cause permanent changes in subsequent generations of progeny. (Supported by NIEHS R03 ES018962)

2111 Comparison of Estrogen Mixtures In Vitro vs In Vivo.


Numerous sources contribute to widespread contamination of drinking water sources with both natural and synthetic estrogens, which is a concern for potential ecological and human health effects. In vitro screening assays are valuable tools for identifying mechanisms of toxicity but in vitro results cannot be directly extrapolated to in vivo exposures since most in vitro assays do not account for metabolism.
distribution and excretion or other systemic toxicities. In this study, we highlight some of the limitations associated with using in vitro estrogen transcriptional activation assays for predicting in vivo action of xenoestrogens. In particular, we compared the ability to predict the uterine growth response (uterotrophic assay, UA) to estrogenics, administered to the rat orally either individually or as mixtures,, using an in vitro estrogen transcriptional activation (TA) assay (T4/TD-hlbicu cell line). We demonstrated that the in vitro activity of the estrogenic agent (BBP) + Methoxychlor in the UA conforms to dose additive (DA) estrogenicity, whereas the degree of estrogenicity of this mixture is underestimated by the TA assay. In contrast, the TA assay responded to a binary mixture of benzenzylphthalate (BBP) + BPAF in a DA manner, whereas, the UA displayed no estrogenic response to this mixture. These data illustrate the limitations associated with making in vivo predictions based on in vitro assay data for compounds that are metabolically inactivated in vivo in the liver, gut or other tissues or activated by the liver in vivo. Ongoing efforts related to this study include characterizing individual dose response curves and mixture estrogenicity for additional estrogens in vitro and in vivo. These data will be used to make predictions from the in vitro assay to the in vivo response to exposure to the compounds. This information is critical for valid interpretation of in vitro screening assay results. Disclaimer: Abstract does not necessarily reflect U.S.EPA policy.

2112 Bisphenol A May Affect the Fertilizing Ability of Mouse Oocytes via Mechanisms Involving Events from Sperm Penetration into the Oocyte to Formation of 1-Cell Zygote.


The cumulus cells surrounding the oocyte expand before ovulation to allow oocyte maturation, and to facilitate sperm cell to penetrate the oocyte. It has been demonstrated that bisphenol A (BPA), a plasticizer that leaches from plastics into food and water, alters the expansion of cumulus cells. Since we have shown that BPA decreases the ability of oocytes, surrounded by cumulus cells, to be fertilized by sperm cells, this study examined whether BPA affects the fertilizing ability of oocytes through effects on cumulus cells, oocyte penetrability by sperm cells or zygote development to 8-cell stage. Female C57BL/6j mice (n = 6-8 per group) were exposed orally to BPA (50 μg/kg/d), diethylstilbestrol (10 μg/kg/d, positive control) or corn oil during 3 estrous cycles every 24 h. Following treatments, mice on estrus received equine chorionic gonadotropin hormone (hCG; 5 IU) ip, 48 h later mice received hormone human chorionic gonadotropin (hCG; 5 IU) and 16 h post-hCG mice were subjected to superovulation and were treated with equine chorionic gonadotropin hormone (hCG; 5 IU). In vitro fertilization (IVF) was carried out on Day 2 post-hCG by insemination of cumulus-free oocytes with spermatozoa from the same strain. In order to evaluate the effect of BPA on fertilization, we determined the percentage of penetrated and fertilized oocytes by staining with acridine orange and rhodamine 123. Data were expressed as % penetrations and % fertilizability. Differences were analyzed by standard t test. BPA at a concentration of 200 μg/kg/d caused no reduction in the percentage of penetrated (Pen) and fertilized (Fert) oocytes compared to vehicle control. In the PND 90 mammary gland, both EE2 doses induced ~ 3x the expression of Esr2, while the expression of Esrrb was induced ~3.5x by the low, 25 μg/kg/d EE2 dose. BPA did not alter the expression of the genes analyzed in control tests. In the mammary gland, the most expressed gene analyzed was ERα while the expression of Esr2, which encodes ERα, and G-protein-coupled ER (Gper) was analyzed in the whole prostate and female mammary gland. The expression of the other receptor genes was 1% Gapdh or less and was not affected by treatment with BPA. In the PND 90 mammary gland, the most expressed gene analyzed was ERα and its expression was also high in adult prostate with no significant changes induced by treatment with BPA. The expression of the other estrogen receptor genes, such as squamous metaplasia, was reduced by two of the binders. The results of this experiment demonstrated how morphometric parameters can be used to assess the effects of xenoestrogens, such as ZEA, and the potentially receptor-dependent ameliorative effects of proprietary binders.

2113 Amphoteric Fluorotelomer-Based Surfactant: 28-Day Subchronic and One-Generation Reproduction Toxicity in Rats.

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An amphoteric fluorotelomer-based surfactant in glycol solvent mixture and water was evaluated in a 28-day oral gavage study (OECD 407) with a 28-day recovery subset and a one-generation reproduction study subset (OECD 422). Groups of 20 Gt/Cd(SD) rats were dosed with vehicle (deionized water) containing 0, 10, 50, or 200 mg/kg/day test substance. Dams were allowed to deliver and rear their offspring until postnatal day (PND) 4. Litter examinations were evaluated at birth and on PND 4 for sex, body weight, and gross postnatal examinations were performed on selected rats and selected organs were weighed and/or retained for histopathological examination. There were no test substance-related deaths or clinical observations, no effects on body weight or nutritional parameters, no effects on neurobehavioral endpoints, clinical pathology, reproductive performance, or on offspring at any dose. Test substance-related changes occurred at ≥ 50 mg/kg/day in the kidneys of male rats and at ≥ 50 mg/kg/day in the cortic tubules of males after 28 days of administration, and were also observed in the P1 male after 45 days of test substance administration. Increased hyaline droplets consistent with α2u globulin were noted at 28 days and/or retained for histopathological examination. There were no test substance re-responses observed in females. Increased hydroxychloroquine in the DA conforms to dose additive (DA) estrogenicity, whereas the degree of estrogenicity of this mixture is underestimated by the TA assay. In contrast, the TA assay responded to a binary mixture of benzenzylphthalate (BBP) + BPAF in a DA manner, whereas, the UA displayed no estrogenic response to this mixture. These data illustrate the limitations associated with making in vivo predictions based on in vitro assay data for compounds that are metabolically inactivated in vivo in the liver, gut or other tissues or activated by the liver in vivo. Ongoing efforts related to this study include characterizing individual dose response curves and mixture estrogenicity for additional estrogens in vitro and in vivo. These data will be used to make predictions from the in vitro assay to the in vivo response to exposure to the compounds. This information is critical for valid interpretation of in vitro screening assay results. Disclaimer: Abstract does not necessarily reflect U.S.EPA policy.

2114 Effects of Zearalenone with or without Proprietary Binders on Vagal Morphology of Prepubertal Gilts.

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The xenoestrogenic mycotoxin, zearalenone (ZEA), can cause hyperestrogenism in prepubertal gilts, resulting in changes in the size, structure, and, potentially, function of the female reproductive tract. Objectives of the present study were to: 1) evaluate ZEA-induced changes in the vaginal epithelium using morphologic parameters and 2) determine whether proprietary binders can ameliorate the xenoestrogenic effects of ZEA. Thirty prepubertal gilts were assigned to five treatment groups. The gilts were housed individually and fed either a control diet, the same control diet with an added 1.5 mg of ZEA per kg of feed, or the ZEA-contaminated diet with Binder A, B, or C added. Animals were humanely sacrificed on Day 21 and portions of the reproductive tracts were removed and fixed in formalin. Image analysis software was used to measure total vaginal lumen circumference, as well as the total length of vaginal epithelium exhibiting hyperplasia and/or squamous metaplasia. The mean vaginal luminal circumference of the cross sections was greater in ZEA-treated groups and the controls, with or without binder. Likewise, the mean percentage of hyperplastic vaginal epithelium was also higher in the ZEA-treated groups, regardless of the presence of binder. There was a tendency for two of the binders to have ameliorative effects on the percentage of vaginal epithelium undergoing ZEA-induced squamous metaplasia. It is clear that treatment with ZEA affected the morphometric parameters evaluated. While none of the binders appeared to ameliorate the effects associated with ZEA, those effects of ZEA and its metabolites involving ER-β estrogen receptors, such as squamous metaplasia, were reduced by two of the binders. The results of this experiment demonstrated how morphometric parameters can be used to assess the effects of xenoestrogens, such as ZEA, and the potentially receptor-dependent ameliorative effects of proprietary binders.

2115 Effect of Bisphenol A (BPA) and Ethinyl Estradiol (EE2) in the Gene Expression of Estrogen Receptors (ER) and ER-Related Receptors in the Rat Prostate and Mammary Gland.


Tissue and temporal expression of receptors involved in estrogen signaling and their modulation by hormonally active agents are hypothesized to influence the long term effects of such agents; however, these expressions are poorly defined. NCTR Sprague-Dawley rats were dosed from gestation day 6 until parturition by oral gavage and their pups were directly dosed by the same route from postnatal (PND) 1 to 90. Dose groups included naïve and vehicle controls, BPA (2.5 μg-300 mg/kg body weight [bw]/day), and EE2 (0.5 and 5.0 μg/kg bw/day). The expression level of genes coding for nuclear ERs (Esr1 and 2), ER-related receptors (Era, E2, and g), and G-protein-coupled ER (Gper) was analyzed in the whole prostate and female mammary gland at PND 4 and 90. Quantitative real-time RT-PCR was used and data was expressed as % Gapdh expression level. The most highly expressed receptor in PND 4 prostate was Era and its expression was also high in adult prostate (3 and 30% Gapdh, respectively). The expression of Era2, which encodes ERβ, was only 0.1% Gapdh in PND 4 prostate, but increased to 40% Gapdh at PND 90. The expression of the other receptor genes was 1% Gapdh or less and was not affected by age (Esr1 > Era2 > Gper > Egrp). Neither BPA nor EE2 affected the expression of the receptor genes analyzed in the prostate under our conditions. In the mammary gland, the trend was the same. The expression of Era2 (0.5% Gapdh) was the least expressed gene was Era2 (0.05% Gapdh), Era2 and Era1 at PND 4 and 90. At PND 4, Era2 was slightly (~<2x) induced by low EE2 dose, relative to vehicle control. In the PND 90 mammary gland, both EE2 doses induced ~3x the expression of Era2, while the expression of Era1 was induced ~3.5x by the low, but not high, EE2 dose. BPA did not alter the expression of the genes analyzed in the mammary gland. Our data suggest that the expression of these receptors is tissue-specific and that BPA and EE2 differentially modulate their expression. IAG FDA 224-12-0003/NIH ES12013.
2116 A Combined Repeated Dose and Reproductive/Developmental Toxicity Screening Study of Perfluoroundecanoic Acid in Rats. 
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Perfluoroalkyl carboxylic acids are one of environmental contaminants which have received attention because of their possible effects on wild life and human health in recent years. In order to obtain the initial risk information on the toxicity of perfluoroundecanoic acid (PFUnA (C11)), we conducted the repeated dose and reproductive/developmental toxicity screening test. PFUnA was administered by gavage to rats at 0 (vehicle: corn oil), 0.1, 0.3 or 1.0 mg/kg/day. Males were dosed for 42 days beginning 14 days before mating and females were dosed from 14 days before mating to day 4 of lactation. During the dosing period, body weight gain was inhibited in both sexes at 1.0 mg/kg/day. In this group, there was a decrease in fibrinogen in both sexes and shortening of activated partial thromboplastin time in males. Blood biochemical examination revealed an increase in BUN and decrease in total protein in both sexes and increases in ALP and ALT and decrease in albumin in males at 1.0 mg/kg/day. The relative liver weight was increased in males at 0.3 mg/kg/day and above and in females at 1.0 mg/kg/day, and histopathologically, centrilobular hypertrophy of hepatocytes was observed in both sexes at 0.3 mg/kg/day and above. Focal necrosis and diffuse vacuolation of hepatocytes were also found in the 1.0 mg/kg/day group. Regarding the reproductive/developmental toxicity, the body weight and pup birth weight and body weight gain for 4 days after birth was inhibited at 1.0 mg/kg/day while no dose-related changes were found in the other reproductive/developmental parameters. Based on these findings, the NOAELs for the repeated dose and reproductive/developmental toxicity are considered to be 0.1 mg/kg/day and 0.3 mg/kg/day, respectively.

2117 The Detrimental Effect of Bisphenol A on Mouse Two-Cell Zygote May Depend on the Dose. 
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Early development of the zygote begins with the first cleavage that forms diploid cells and it depends on the contribution of paternal gametes. Bisphenol A (BPA) is an environmental compound that may leach from plastics into food or water. Some studies have shown that BPA alters embryonic development but they mostly focus on effects at later stages of development. This study evaluated the effect of the in vivo exposure to different doses of BPA on the formation of two-cell zygotes. Female mice C57BL/6J (30 days old) (n = 6-16 per group) were orally exposed to BPA (0.5, 5 and 50 mg/kg/d) or corn oil (isophorone-stripped, negative control) for 3 estrous cycles every 24 h. Following treatments, mice on estrus received equine chorionic gonadotropin hormone (5 IU) ip, 48 h later mice received hormone human chorionic gonadotropin (hCG; 5 IU) and 16 h post-hCG mice were bred with a fertility proven male mouse to collect two-cell zygotes. Zygotes were stained with Hoechst 33342 and classified for abnormalities such as cell lysis and cytoplasmic prolongations. Mice treated with all doses of BPA had lower percent of fertilized oocytes compared to control, but zygote abnormalities were observed from the dose of 0.5 mg/kg/d BPA. Specifically, two-cell zygotes from mice exposed to 0.5 mg/kg/d BPA had higher percent of cell lysis or cytoplasmic prolongations compared to control, and two-cell zygotes from mice exposed to 50 mg/kg/d BPA had higher cytoplasmic prolongations compared to control. Our data suggest that BPA alters the formation of 2-cell zygote causing abnormalities depending on the dose. Conacyt-Mexico CB-167678.

2118 Body Burden and Preliminary Effects in Rats following Low-Dose Drinking Water Exposure to a VOC Mixture during Pregnancy and Adolescence. 
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Volatile organic compounds (VOCs) such as benzene, trichloroethylene, trans-1,2-dichloroethylene, tetrachloroethylene, and vinyl chloride are common industrial solvents used in a number of cleaning agents and solvents. Ingestion or inhalation of these compounds are the most common routes of reported environmental or occupational exposures to VOCs. In the US, in births defects, infant mortality, reproductive cancers and leukemia have occurred in areas where high levels of VOCs were detected in drinking water supplies. Here, time-pregnant Harlan Sprague Dawley rats and their offspring were given access to water containing mixtures of these 5 VOCs at concentrations 5, 10 and 50 times those detected in contaminated US drinking water (Sonnenfeld et al., 2001). Dams and pups were exposed from gestation day (GD) 10 until sacrifice. Blood was collected under hermetic conditions to determine VOC body burden in both dams (GD13, 15, 20 and postnatal day (PND)15, 21) and pups (PND15, 21, 28, 48). In exposed dams, low levels of VOCs were detected in the blood during pregnancy and increased during lactation. Dose dependent changes in each compound was also evident. Blood levels did not plateau, and differences in VOC body burden were attributed to the varying amounts of water consumed. Pup VOC levels also varied with changing body weight, milk and water consumption. At necropsy, pup body, liver and spleen weights were recorded and various tissues were collected for further analysis. VOC-exposure had no effect on selected organ weights or body/organ weight ratios. Morphological changes in the mammary gland were detected and were most prominent in male mammary tissue. Disclaimer: This abstract does not necessarily reflect NIEHS and CDC policy.

2119 The Inhibiting B (InhB) Response to the Testicular Toxics Mono-2-Ethylhexyl Phthalate (MEHP), 1,3-Dinitrobenzene (DNB) or Carbendazim (CBZ) Following Short-Term Repeat Dosing in the Male Rat. 
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The objectives of this study were to evaluate the utility of plasma InhB as a biomarker of testicular injury in adult rats using the known Sertoli cell toxicants MEHP, DNB or CBZ. The studies were run under conditions of short-term repeat dosing similar to that which would be used in early stage drug development to screen and prioritize drug candidates. The short-term high-dose exposure paradigm also allowed for assessment of changes in InhB as an early indicator of testicular injury. Following oral gavage administration of the compounds for 2 or 7 days, the rats were evaluated for clinical signs, body weight, food consumption, organ weights, plasma hormone levels, and gross and microscopic pathology of selected organs. MEHP and DNB, and CBZ were found to be characterized by minimal exfoliation of germ cells as demonstrated by increased cellular debris in the epididymis (MEHP) to more severe and dose/duration responsive Sertoli cell vacuolation, germ cell degeneration, and multinucleated giant cells of germ cell origin (DNB and CBZ). The slight to moderate Sertoli and germinal cell injuries did not correlate with significant changes in plasma InhB levels following 2 or 7 days of exposure. However, moderate to severe injury to germinal epithelium following up to 7 days of exposure, but not after a 2 day exposure, correlated with decreased in plasma InhB levels and less consistently with increases in plasma follicle stimulating hormone (FSH). In conclusion, under the conditions of these studies, changes in InhB were not an effective early onset marker of testicular toxicity or an effective marker for slight to moderate levels of acute injury and only reflected more severe disruption of spermatogenesis. Changes in plasma InhB and FSH were poorly correlated except in some instances of moderate to marked testicular toxicity.

2120 Poor Correlation between Rat Testis Histology and Serum Inhibin B after Treatment with Two Drug Candidates. 
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Serum InhB was measured in two studies of known-testis-toxic drug candidates. Study 1 was for a Hepatitis C candidate, and utilized a 10 week dosing period, followed by mating and necropsy of half of each group, and then a 12 week recovery period for the remaining 15 rats/group. At the end of dosing-post mating necropsy, 6 of 15 high-dose males had testis lesions (germ cell loss, degeneration); InhB was significantly reduced in all animals in that group. The mid-dose group had no testis lesions but significantly reduced mean serum Inhibin B. After the 12 week recovery, 9/15 high-dose males showed damage in testes, and no mid-dose animals had testis lesions. Mean serum Inhibin B in all treated groups at recovery was not different from controls. Inhibin B appeared to both over-report and under-report testis damage in Study 1. Study 2 was an acute pathogenesis study for an antibacterial compound, using control and two dose levels and multiple time-points (days 5, 8, 15, 22, and then untreated until day 71). At each timepoint, blood was sampled from all remaining rats and 5/group were killed for histologic evaluation. The low-dose group developed minimal to moderate lesions, while
serum Inhibin B was never changed. The high-dose animals progressed quickly from (mainly) spermatogenic to being broadly and moderately affected and finally died (cell absence, disorganization); serum Inhibin B levels were reduced at days 8 and 15 only. In this study, Inhibin B appeared less sensitive than histology, except with marked testis damage, when Inhibin B was routinely low. Serum Inhibin B both over-reported damage (being reduced in the absence of lesions) and under-reported testis damage (being normal in the presence of testis lesions) in these two studies. We conclude that across both of these studies, there was a poor correlation between changes in serum levels of Inhibin B and testis histopathology.

2121 Assessment of Inhibin B As a Biomarker of Testicular Injury following Administration of Carbendazim, Cetrorelix, or 1, 2-Dibromo-3-Chloropropane in Wistar Han Rats.
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Although histopathology is considered the gold standard for assessing testicular toxicity in the nonclinical setting, identification of non-invasive biomarkers for testicular injury are necessary to improve safety monitoring capabilities for clinical trials. Inhibin B has been investigated as a potential noninvasive biomarker for testicular toxicity. The present study investigates the correlation of Inhibin B serum levels in Wistar Han rats with the onset and reversibility of testicular histopathology from classical testicular toxicants, carbendazim (CBZ), cetrorelix acetate (CTX), and 1,2-dibromo-3-chloropropane (DBCP). The dosing paradigm was selected with the intent of monitoring the impact of each toxicant on the testis from the onset of testicular pathology occurring at any additional dosing in the Recovery Phase, only assessing a leading indicator of the onset of testicular toxicity, prior to the onset of germ cell depletion. However, since Inhibin B was not increased at the end of the Dosing Phase and the onset of testicular pathology occurred without any additional dosing in the Recovery Phase, it is clear that monitoring Inhibin B would provide sufficient advanced warning of the onset of testicular pathology. Furthermore, FSH was increased and the ratio of Inhibin B/FSH was increased with DBCP administration in the Interim Phase, but not in the Drug or Recovery Phases. Although the Inhibin B/FSH ratio was a leading indicator of testicular pathology, the effective window for monitoring may be narrow. Conclusion: Inhibin B has limited predictive capacity as a leading testicular biomarker in rats.

2122 The Inhibin B Response in Male Rats Treated with a GnRH Agonist and an Endothelin Receptor Antagonist.
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The tests show a moderate frequency as a preclinical toxicity target organ. This is primarily detected by histopathology and there is a need to identify circulating biomarkers to enable longitudinal monitoring and facilitate safe progression of compounds into the clinic. Inhibin B is primarily synthesized by the Sertoli cells and regulates pituitary FSH release through a negative feedback loop. This study was part of a HESI-sponsored initiative to evaluate Inhibin B as a marker of spermatogenic dysfunction in the rat. Inhibin B was measured in male Han Wistar rats (10 weeks old) administered vehicle or an endothelin receptor antagonist (ET-An) orally for 28 days or a GnRH agonist (GnRH-A) as a subcutaneous implant on Day 1. Ten animals/group/time point were killed on Days 4, 8, 15 and 29 (controls on Days 15 and 29), for testes weights and histopathology. In-life blood samples were taken on Days 4, 8, 15 and 29 to measure inhibin B, FSH and LH, and at necropsy for the same hormones plus testosterone. Plasma inhibin B showed a wide concentration range in control animals (group means 7.64 to 18.42 pg/mL; individual animals 17.8 to 381 pg/mL). GnRH-A caused decreased testes weights plus degenerative testicular pathology from Day 4 with partial recovery by Day 29. Statistically significant reductions in inhibin B were observed at all time points and appeared to track the development and partial recovery of the pathology (generally <50 pg/mL on Days 4 to 15; group mean 92 pg/mL on Day 29). ET-An produced an increase in testes weights and a non degenerative lesion of minimal tubular dilatation. There was a trend for lower inhibin B values (30 to 50%) at all time points, including on Day 4 when tubular dilatation was not yet evident. Overall, we conclude that following GnRH-A administration, inhibin B showed a good correlation with testicular pathology for GnRH-A, and following ET-An administration appeared to give a signal that might reflect changes in tubular function in the absence of degenerative pathology.

2123 Inhibin B As a Marker of Sertoli Cell Damage and Spermatogenic Disturbance in the Rat.
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This study was designed to determine the effects of Compound A on the fertility and early embryonic development in the male rat over a 15-19 weeks treatment and a 19 weeks treatment free period in control and 30, 60 and 180 mg/kg/dose groups (n=22/group). Compound A is a dose-dependent manner induced various degrees of spermatogenic alterations compatible with Sertoli cells being the primary target, e.g. inter- and intracellular Sertoli cell vacuolization and altered cellular morphology followed by germ cell degeneration and marked reduction of epididymal sperm numbers. Blood-testis barrier remained intact (electron microscopy and hyperosmotic fixation test) until germ cells disappeared. Mating behaviour and weights of androgen-dependent prostate and seminal vesicles remained unaffected. Inhibin B levels correlated only with moderate to severe spermatogenic alterations. Ten animals with inhibin B levels below detection limit were encountered and five of these animals were fertile in week 19 but following another 15 weeks without treatment, animals were rendered infertile and inhibin B levels remained undetectable. In the rat, inhibin B only reflects major spermatogenic alterations and markedly reduced inhibin B levels might indicate irreversibility of these alterations and even infertility.

2124 The Inhibin B Response to Testicular Toxicants Ethylene Glycol Monomethyl Ether or Dibromoacetic Acid in Male Rats.
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This study was conducted as part of an ILSI-HESI consortium effort to assess the utility of circulating inhibin B as an early biomarker of testicular toxicity in rats. Two known testicular toxicants were selected for use in this study: ethylene glycol monomethyl ether (EGME) and dibromoacetic acid (DBAA). EGME (200 mg/kg/day), DBAA (250 mg/kg/day) or vehicle control (0.2% hydroxypropyl methylcellulose [HPMC]) were administered orally to male rats for 3, 6 or 14 consecutive days. On study days 4, 7, and 15, serum was collected for evaluation of inhibin B levels from all surviving animals and subset of animals was necropsied from each of the control, EGME and DBAA groups. Administration of EGME resulted in spermatocyte degeneration in late stage tubules and spermatocyte depletion to stage III on day 4, progressing to loss of spermatocytes and round spermatids to stage VI by day 7 and continued germ cell loss and degeneration of elongating spermatids by day 15. Inhibin B levels among EGME-treated animals progressively decreased relative to their respective controls at all time points. Administration of DBAA was associated with spermatid retention at all three time points and abnormal residual bodies at days 7 and 15. Inhibin B levels among DBAA-treated animals decreased progressively relative to their respective controls on days 7 and 15. The results of this study indicated that serum inhibin B levels in rats provided a signal of testicular toxicity for each of these known testicular toxicants administered at high levels; however, histopathology provided the earliest evidence of toxic effects.

2125 The Inhibin B Response to the Testicular Toxin, 1, 3-Dinitrobenezene in Rats and the Analytical Evaluation of Inhibin B ELISA Kit.
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Background: This work is part of an ILSI-HESI consortium effort to evaluate the analytical performance of a second generation ELISA kit and also to assess the utility of circulating inhibin B (InhB) as an early biomarker of testicular toxicity in rats. Methods: A commercially available InhB ELISA was used to assess for dilution linearity, frozen stability and serum/plasma comparison. Reference ranges were generated for male Sprague Dawley rats. To evaluate the biological utility of InhB, 1, 3-dinitrobenezene (DNB), a Sertoli cell toxicant, was orally administered to male rats for 2 or 5 consecutive days at 2 or 6 mg/kg/day. On Days 1 and 2, serum was collected for evaluation of InhB and all rats were treated for 2 days, and on Days 1, 3 and 5 from all rats treated for 5 days. At the end of treatment, testes were weighed and examined histologically. Results: There...
was no difference between serum or plasma InhB values and they were stable out to 12 weeks when stored at -20°C and -80°C. Dilution linearity was acceptable up to 32-fold. An age-related decline of InhB levels was seen between 6 and 9 weeks of age after which levels were stable up to 20 weeks. DNB caused a time-dependent increase in incidence and severity of testicular findings characterized by degeneration of the germinal epithelium, loss of pachytene spermatocytes and vacuolization of the Sertoli cells. Decabromodiphenyl ether (decaBDE) caused a high dose effect on Day 15 and without any associated changes in FSH. Conclusions: Overall, the InhB assay performed well under our conditions; however it is important to be aware of the biological variability and low control values observed by some other laboratories. In our study, a change in serum InhB levels was detected only in association with moderate/severe testicular toxicity, and is therefore considered of limited value as an early biomarker for Sertoli cell toxicity.

### 2126 The Inhibin B (InhB) Response to the Testicular Toxicants Hexachlorophene, Ethane Dimethane Sulfonate (EDS), Dibutyolphthalate (DBP), Nitrofurazone, DI-Ethionine, 17-Alph Ethyleneslradiol, 2, 5-Hexanediene, or Carbendazim (CBZ) following Short-Term Dosing in the Male Rat.

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Background: InhB is a hetero-dimer glycoprotein that down regulates follicle stimulating hormone and is produced predominantly by the Sertoli cells. The potential correlation between changes in plasma InhB and Sertoli cell toxicity was evaluated in male rats administered various testicular toxicants in 8 separate studies. InhB fluctuations over 24 hours were also measured. Methods: For the testicular toxicity studies, five to eight Sprague-Dawley, Wistar, or Wistar-Han rats ranging from 8 to 13 weeks of age were administered 1 of 8 testicular toxicants for 1 to 29 days (depending on the compound). The 8 testicular toxicants were Dl-Ethionine, dibutyolphthalate, nitrofurazone, 2,5-hexanediene, 17-alpha ethyleneslradiol, ethane dimethane sulfonate, hexachlorophene, and carbendazim. For the 24-hour time period, plasma was collected by an automatic blood sampler. Results: Histomorphologic testicular findings were seminiferous tubule degeneration (STD), apoptosis and pachytene spermatocytes and vacuolization of the Sertoli cells. Across the 8 testicular toxicants tested there were varying degrees of correlation between decreases in InhB and STD and spermatogenesis. In an ROC exclusion model analysis, where treated samples without histopathology were excluded, performed on all studies except EDS (Leydig cell toxicant), InhB showed a sensitivity of 72% at 90% specificity, demonstrating the potential value of InhB as a biomarker of testicular toxicity. Conclusion: Decreases in InhB showed a good correlation with Sertoli cell toxicity. As anticipated, there was no correlation between decreases in InhB and Leydig cell toxicity (interstitial cell degeneration). A pattern of InhB secretion could not be identified over a 24 hour time period.

### 2127 Simvastatin and Dipentyl Phthalate Lower Testosterone Production and Exhibit Dose Additive Effects on the Fetal Testis via Distinct Mechanistic Pathways.

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Sex differentiation of the mammalian reproductive tract is a highly regulated process that is driven, in part, by fetal testosterone (T) production. In utero exposure to phthalate esters (PE) during sex differentiation can result in reproductive tract malformations in rats. PE alter the expression of genes associated with steroid synthesis/transport and cholesterol biosynthesis. Simvastatin (SMV) is a cholesterol-lowering drug that interferes with cholesterol biosynthesis. As cholesterol is a precursor for steroid biosynthesis, we proposed that like PE, maternal exposure to SMV during the critical period of sex differentiation would lower fetal T production and ultimately result in female reproductive tract malformations. Simvastatin (SMV) is a cholesterol-lowering drug that interferes with cholesterol biosynthesis. As cholesterol is a precursor for steroid biosynthesis, we proposed that like PE, maternal exposure to SMV during the critical period of sex differentiation would lower fetal T production and ultimately result in female reproductive tract malformations. DPeP treatment group in the form of malocclusions and incomplete zygomatic ossification. These results indicate the DPeP is about 3.5 fold more potent in inducing the PS in F1 male rats than is DEHP and demonstrate that the relative potencies for disrupting fetal testis endocrine function can be used to predict some of the postnatal reproductive effects of this class of endocrine disruptors. Disclaimer: This abstract doesn’t necessarily reflect USEPA policy. Supported in part by NTP/NIEHS I# RW-75-92285501.

### 2128 Postnatal Effects of Dipentyl Phthalate on Male Rat Reproductive Development.

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We conducted several in utero, ex vivo and in vitro studies to characterize the relative potencies of a series of phthalates on fetal rat testis testosterone production and gene expression. Dipentyl phthalate (DPeP) was the most potent of the active chemicals in its effect on fetal testis endocrine function. Although these studies have pointed to the overall potency of DPeP, little literature exists defining its dose-response curve in vivo. The objective of this study was to determine if the potency of DPeP on fetal testis endocrine function was predictive of the ability of the chemical to induce reproductive tract malformations in male rats. Male offspring of treated dams displayed decreased AGD, increased nipple retention, incomplete preputial separation, decreased sperm production, hypospadias, undescended testicles, ormalformations of the testes, ventral prostate, and seminal vesicles, reduced body weight (300 mg/kg/day) and reduced postnatal survival. Phthalate syndrome (PS) malformations were seen in about 9%, 44% and 100% of the F1 male offspring at 33, 100 and 300 mg/kg/d, dosage levels that reduced fetal T production by 5%, 35% and 93% respectively. Also of note were skull malformations in highest treatment group in the form of malformations and incomplete zygomatic ossification. These results indicate the DPeP is about 3.5 fold more potent in inducing the PS in F1 male rats than is DEHP and demonstrate that the relative potencies for disrupting fetal testis endocrine function can be used to predict some of the postnatal reproductive effects of this class of endocrine disruptors. Disclaimer: This abstract doesn’t necessarily reflect USEPA policy. Supported in part by NTP/NIEHS I# RW-75-92285501.

### 2129 Changes of Expression Levels of Oxidative Stress-Related Genes in Mouse Epididymides by Neonatal Exposure to Low-Dose Decabromodiphenyl Ether.

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Decabromodiphenyl ether (decaBDE), one of polybrominated diphenyl ethers (PBDEs), is the most famous flame retardant and is used worldwide. In a previous study, we identified adverse effects of neonatal decaBDE exposure on mouse epididymides, for example decrease of epididymal weight. On the other hand, neonatal exposure to diethylstilbestrol (DES), artificial estrogenic compounds, also causes several adverse effects on epididymides. DES exposure causes the decrease of epididymal weight, morphological abnormality, and the lasting change of expression levels of several genes. Molecular mechanisms for induction of harmful effects by decaBDE exposure remain unclear. Since many studies have reported that PBDEs have estrogenic activity, this activity may contribute to induction of adverse effects of decaBDE exposure. This study was carried out to examine how effects caused in epididymides by neonatal decaBDE exposure, and elucidate molecular mechanisms of adverse effects by decaBDE exposure. Administration of decaBDE was performed subcutaneously at 0.25 mg/kg body weight/day, on postnatal days 1 to 5. At 12 weeks of age, epididymides were histologically examined and gene expression was analyzed using DNA microarray and real-time PCR. Our data showed that 1) no histological change was observed on epididymal tissues by neonatal decaBDE exposure, differently from that of DES, 2) decaBDE exposure could not induce the changes of expression levels of genes which were affected by DES, but caused changes of expression levels of some oxidative stress related genes in mouse epididymides, 3) the expression level of Ubiquitin C (Ubc) increased in decaBDE-exposed mouse epididymides. Our presented data suggest the possibility that increase of oxidative stress is related to elicitation of harmful effects in decaBDE-exposed mouse epididymides.
- Wistar male (male) were divided into two groups — group A (GA) served as control and group B (GB) were daily administered orally chlorpyrifos (Anu Products Ltd., India) at a dose of 5 mg/kg b.wt. Rats were sacrificed on 1st, 2nd, 4th, 6th, and 8th week after initiation of the experiment. Left testes and duodenum were extracted and fixed in aqueous Bouin's solution. The tissues thus fixed were routinely processed for histological studies. Chlorpyrifos treatment caused degeneration in seminiferous tubules and thus inhibit the spermatogenesis in rats. Few seminiferous tubules lack the germinal epithelium. After exposure to chlorpyrifos the morphological observations of duodenum showed an increased mucous cell activity, disruption and sloughing of duodenal villar cells, lymphocytic infiltration and degeneration of cells. In conclusion the findings of the present study indicate that the organophosphate — chlorpyrifos can inhibit spermatogenesis and provoke degenerative features in the duodenum of rats and thus severely affect these organs.

Phthalate esters (PE) such as diethylhexyl phthalate (DEHP) produce reproductive malformations in male rodents by reduction of testosterone (T) production and gene expression after dams are exposed during the critical period of sexual differentiation. We investigated the effects of seven PE known to reduce fetal T production and further evaluated gene expression using novel SABiosciences PCR arrays. Each array tests for 84 genes either involved in phase one or drug lipoprotein transport and cholesterol metabolism/synthesis. Timed pregnant Sprague-Dawley rats were orally dosed with individual phthalates from gestational day (GD) 14-18. On GD 18, testes were collected from 3 fetuses per dam and cultured for 3 hours. Medium was collected and T values measured by RIA. Remaining testes were pooled by litter, RNA extracted, cleaned, quantified, and evaluated for gene expression using PCR arrays. PE were DiHP (750 mg/kg), DiHexyl (750 mg/kg), DPP (100 mg/kg), DBP (900 mg/kg), DNS (1500 mg/kg), DCHP (0, 100, 300, 600, or 900 mg/kg), and DEHP (750 mg/kg). Of the 84 genes on the Drug Metabolism arrays, four were significantly reduced: Cyp11b1, Cyp11a1, Cyp17a1, and ALDH2. Cyp11a1 and Cyp17a1 are involved in the production of cortisol and corticosterone. Using the lipoprotein/cholesterol arrays, we observed that a mixture of 9 PE administered on gestation day (GD) 14-18 at doses of 11, 33, 100, or 300 mg/kg/d (n=3); controls received the vehicle corn oil. At GD 18 dams were necropsied and fetal specimens (plasma, testes, repro) were collected and T values measured by RIA. Remaining testes were pooled by litter, RNA extracted, cleaned, quantified, and evaluated for gene expression using PCR arrays. PE were DiHP (750 mg/kg), DiHexyl (750 mg/kg), DPP (100 mg/kg), DBP (900 mg/kg), DNS (1500 mg/kg), DCHP (0, 100, 300, 600, or 900 mg/kg), and DEHP (750 mg/kg). Of the 84 genes on the Drug Metabolism arrays, four were significantly reduced: Cyp11b1, Cyp11a1, Cyp17a1, and ALDH2. Cyp11a1 and Cyp17a1 are involved in the production of cortisol and corticosterone. Using the lipoprotein/cholesterol arrays, we identified 10 genes that were significantly altered that are involved in cholesterol biosynthesis (such as APOC3, Hdc2c67, EBP, MVK, Tm7s2). These data support that PE may alter alterations in steroidogenesis and include genes for other hormones (Insl3), growth factors, steroid transport proteins, and multiple genes involved in cholesterol synthesis. Disclaimer: This abstract does not reflect USEPA policy. Supported by NTP/NIEHS IA# RW-75-92285501-1 and fellowship administered jointly between EPA/DOE and the Oak Ridge Institute for Science and Education.
reduced fetal T levels and the resultant postmale reproductive malformations. 

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2135 Impact of Clinically Relevant Cisplatin Treatment on the Undifferentiated Spermatogonia Population and Niche.

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A typical clinical cisplatin (CDDP) regimen consists of repeated cycles of 5-7 daily low dose injections followed 1-2 week recovery period; and while effective, often results in prolonged, sometimes permanent, infertility. Theoretically, undifferentiated spermatogonia (undiff-Sp), including spermatogonial stem cells (SSCs), should repopulate the testis after exposure has ceased. We hypothesize that SCC mitotic activity increases following the initial exposure to CDDP, rendering SSCs increasingly susceptible to CDDP-induced injury during subsequent cycles. We examined changes in the adult C57 undiff-Sp population and niche at days 1, 8, and 16 of the recovery period following a clinically-relevant course of 1 cycle (2.5 or 5.0 mg/kg/d) and 2 cycles (2.5 mg/kg/d) of intraperitoneal CDDP. Histological examination of the testis epithelium showed an increase in damage correlating with exposure dose and cycle number. Apoptosis was measured via TUNEL and found to increase during the recovery period following 1 cycle of 2.5mg/kg and 5.0mg/kg (300%), as well as 2 cycles of 2.5mg/kg 600%) CDDP compared to controls. Immunohistochemistry (IHC) was performed using antibodies specific for FOXO1 (undiff-Sp marker) and GDNF (critical SSC niche protein secreted by Sertoli cells). Analysis of FOXO1 showed a reduction in undiff-Sp during the recovery period following 1 cycle of 2.5 (50%) and 5.0mg/kg CDDP (60%), as well as 2 cycles of 2.5mg/kg CDDP compared with controls (50%); followed by a return to numbers equal with or surpassing controls at 16d recovery. IHC analysis of GDNF revealed a general increase in expression, prominently along the basal membrane, during the recovery period in all treatment groups. These data suggest that a greater dose of CDDP in a single cycle of exposure causes a greater impact on the functional stem cell pool and its niche and multiple cycles of exposure result in a still greater impact than an equivalent cumulative dose. Future experiments focus on whether CDDP exposure targets the SSCs, the Sertoli cells, or both.

2136 Using a Rat Primary Epididymal Cell Model to Evaluate Drug-Induced Inflammation and Granulomas.

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Epididymal inflammation and sperm granuloma formation have been detected in vivo in Sprague Dawley (SD) rats following various treatments. The objective of this in vivo primary rat cell model is to identify compounds that cause epididymal inflammation and granulomas using minimal compound and fewer animals. We had previously shown that compounds produced large increases in transcript levels for the IL6 and GRO cytokines (3 to 10-fold increases), while less potent and negative compounds produced smaller (less than or equal to 2X) or no increases. We evaluated the marketed phosphodiesterase type 4 inhibitors, ibudilast (I), roflumilast (R), and the active metabolite roflumilast N-oxide (RNO), which are known to cause sperm granulomas in rats. Primary cultures of mixed epididymal epithelial cells were isolated and plated on Matrigel 24-well plates and exposed to increasing concentrations of test compounds. 24 hours post dosing, we measured mitochondrial metabolic activity using the MTS assay and cell lysates were evaluated for IL6, GRO, and IL10 gene expression by qRT-PCR at the IC20 (80% MTS activity). R and RNO were poorly soluble, which limited the in vitro exposure levels. Perhaps for this reason, modest increases in IL-6 and GRO were seen with R and RNO treatment (1.2- and 1-fold for R, 1.6- and 1.9-fold for RNO) while MTS activity was not reduced. However, treatment with ibudilast resulted in larger increases in IL6 and GRO, 2.7- and 2.9-fold respectively at the IC20, consistent with its in vivo activity. Our results of this study demonstrate that this primary epididymal culture model detects inflammatory signals that reflect the in vivo activity of at least one of these compounds in rats.

2137 Evaluation of an In Vitro Model of Spermatogenesis for Predicting Testicular Toxicity.

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Due to the complex physiology of the testes, in vitro models have been largely unsuccessful at predicting testicular toxicity in vivo. These models often bear limited resemblance to in vivo cellular organization. We evaluated an in vitro model (HuCe et al 1998) that forms tight junctions similar to the blood-testis-barrier (BTB), and supports spermatogenesis through meiosis II (secondary spermatocytes) and the formation of round spermatids. We used this in vitro model to evaluate the toxicity of four known testicular toxicants: Bisphenol A (BPA), 2-Methoxyacetic acid (MAA) 1,3-Dinitrobenzene (DNB), and Lindane in pre-pubertal rat seminiferous tubule cultures treated with compound for 28 days. Formation of the BTB (tight junctions) was measured by transepithelial electrical resistance (TEER) every 2 days. Cells were collected weekly for flow cytometric analysis to measure total viability and cell counts for each stage of spermatogenesis. Concentrations for each chemical were selected to approximate those known to produce in vitro testicular effects. BPA and DNB decreased dramatically the TEER in a dose and time-dependent manner, whereas the effects of MAA and Lindane were less marked and transitory, even at the higher concentrations tested. Cell viability was slightly, if any, modified by the compounds; this was due most likely to the phagocytic activity of the Sertoli cells. All compounds induced dramatic dose- dependent diminutions of the populations of spermatocytes I and II, and round spermatids. It is important to note that the well-known specific toxic effects of MAA and DNB on pachytene spermatocytes were easily reproduced by this in vitro model. Combined, these assays not only predicted testicular toxicity in vitro, but also identified whether the compound directly targets the Sertoli cells or germ cells. Further validation, this cell model may be applicable as a screen to minimize running long-term live-phase studies.

2138 Vinclozolin Alters the Testosterone Homeostasis by Regulating the Rat Liver Cytochrome P450.

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The Vinclozolin (V) is a well-characterized anti-androgenic fungicide in several species. The V is able to produce a complex pattern of induction, suppression and inhibition of cytochrome P450 (CYP) isozymes in mammals. The alteration of liver CYP expression can modify the testosterone biotransformation pattern. The objective of this work was to assess the effect of V on liver CYP expression and its repercussion on serum and urine levels of testosterone and its metabolites. Male Wistar adult rats were orally administered 100 mg/kg/d V for 7 d suspended in corn oil. After last dose the urine of 24 h was collected and animals were sacrificed and the blood was obtained by cardiac punction. The liver was processed to obtain microsomes for enzyme assays and protein content analysis of different CYP isozymes. The testosterone and its metabolites were extracted from plasma and urine and was processed by enzymatic hydrolysis by β-glucuronidase/sulfatase and analyzed by HPLC. V exposure increased 25% the liver relative weight and was accompanied by an increase of 40.3% of total CYP content. The liver immunoreactive protein content of CYP1A1, 1A2, 2A, 2B1 and 3A2 increased 6.5-, 16.4-, 2.3-, 6.5- and 1.5-fold, respect to the non-treated group. The protein content of CYP2E1 was not affected by V. Enzyme activities of EROD, MROD, PROD and PNPH significantly increased 3.0-, 6.4-, 63.3- and 1.6-fold, respectively. In V-treated animals testosteron levels increased 80-fold in serum and 150-fold in urine. V affected the testosterone biotransformation pattern, androstenedione levels increased in serum and urine 54- and 3.6-fold, respectively; 6-DHT increased 112-fold in urine. Other testosterone metabolites were not affected. These results indicate that V regulates liver CYP expression and alters the testosterone biotransformation pattern. In addition, they also suggest an alteration on testosterone homeostasis which may represent another mechanism of action for V. These results will provide further insight into the relationship between toxicity and V exposure.

2139 Sperm mRNAs Are Molecular Markers of Testicular Injury in Rats.

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Traditional endpoints used to measure male reproductive toxicity in humans, including serum and hormone analyses, are insensitive and unreliable; those used to monitor toxicity in animal studies, while sensitive, are not easily translatable to humans. It is therefore necessary to develop sensitive and reliable molecular biomarkers of testicular injury that can be used to both monitor human reproductive function and compare animal studies with human exposures.

Objective: This sub-chronic dose response study aimed to build on existing data that identified 12 mRNA transcripts that were altered in the sperm following exposure to single doses of the Sertoli cell toxicants 2,5-hexadiene (2,5-HD) or carbendazim (CBZ).
Methods: Adult male Fisher 344 rats were either exposed to 0, 0.14%, 0.21%, or 0.33% 2.5-HD in the drinking water for three months, or exposed by daily oral gavage to 0, 30, 50, or 70 mg/kg CBZ in corn oil for three months. Body and organ weights were obtained, and quantifiable histopathological parameters (homogenization resistant spermatid head counts and retained spermatid head counts) were measured. Sperm mRNA were measured by qRT-PCR arrays.

Results: At doses that produced no-to-low levels of histopathological injury, as assessed by organ weights and histopathology, a total of eight mRNA transcripts were altered in the cauda epidymal sperm. Four of the transcripts were significantly increased at the highest dose of HD, with clustin increased in a dose responsive manner at all doses of HD. Six of the transcripts were significantly altered after exposure to CBZ, with clustin also increased in a dose responsive manner at all doses of CBZ.

Conclusions: Our data indicate that sperm mRNA transcripts are sensitive markers of testicular toxicity. The clustin transcript in particular may be a more sensitive indicator of Sertoli cell injury than the most sensitive histopathological endpoint.

2140 Effect of Neonatal Exposure to Decabromodiphenyl Ether on Transcript Levels of Splicing Factors in Mouse Testes.

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Decabrominated diphenyl ether (decaBDE) is one of flame retardants and is used in worldwide. Our previous study found postnatal exposure to decaBDE had adverse effect to male reproduction in mice. Mice were injected subcutaneously 0.025, 0.25, and 2.5 mg/kg decaBDE during postnatal day 1-5 and evaluated at 12 weeks of age. In 0.025, and 0.25 mg/kg dose groups, the number of Sertoli cells was reduced significantly. Thyroid hormones (THs) are key players for Sertoli cells development. We identified that the transcript levels of TH receptor α (Thra) decreased significantly in testes of 0.025 mg/kg decaBDE dose group, while no change of TH levels was observed between control and all dose groups. Moreover, we found the ratio of Thra1/Thra2 level decreased in 0.025 and 0.25 mg/kg dose groups compared to control group. The ratio of Thra1/Thra2 level is known to be varied by the change physiologically, and the ratio of heterogeneous ribonucleoprotein (hnRNPs) A1 (hnRNP1): serine/arginine-rich splicing factor 1 (Srsf1) is related to Thra1/Thra2 ratio level. However, no change was observed in hnRNP1: Smf1 level between control and dose groups. Therefore, we hypothesized other splicing factors may be involved in decrease of Thra1/Thra2 level. In this study, the transcript levels of hnRNPs F (hnRNP1) and hnRNPs H1 (hnRNP1) were examined by real-time PCR. Mice were dosed by above described method, and testes were collected at 12 weeks of age. Our data showed the decrease in transcript level of hnRNP1 in 0.025 mg/kg dose group, and decrease of hnRNP1 level in 0.25 and 2.5 mg/kg dose groups compared to the control group. Our study suggests that hnRNP1 and hnRNP1 are also involved in the decrease of Thra1/Thra2 ratio by neonatal decaBDE exposure.

2141 RSPOs Counteract TCDD Inhibition of Canonical Wnt Signaling during Fetal Mouse Prostate Development.

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Prostatic buds are derived from the urogenital sinus (UGS) and later form into the prostate ductal network in adult mammals. In utero TCDD exposure causes mispositioning and reductive in dorsal and lateral prostatic bud numbers and prevents formation of ventral buds leading to ventral prostate agenesis. Here we examined canonical Wnt signaling following TCDD exposure in vitro. We found multiple components of the pathway (Lef1, Tcl1, Wnt4, Lgr5) to be downregulated by TCDD. R-spondins (RSPOs) are promoters of canonical Wnt signaling but their mechanism for activating the canonical Wnt pathway is not fully understood. Previously, we demonstrated RSPO2 and RSPO3 promote prostatic bud development and addition of these proteins can rescue the effects of TCDD on prostatic bud growth. We now further examined the mechanism of RSPO activation of the Wnt pathway in the UGS by studying the extracellular Wnt antagonists, DKKs, and determining their effects on budding in vitro and examining LGRs, which are putative RSPO receptors, by studying mRNA expression between vehicle and TCDD treated UGSs. Both mechanisms of Wnt activation by RSPOs have been demonstrated in other systems; however, it remains unclear if RSPOs preferentially act through one mechanism over the other. Our results showed little effect on prostate bud number following treatment with DKK1 and DKK2. We found both Lgr4 and Lgr5 mRNA expression in the basal epithelium (BE) where prostotic bud formation and additionally we found a decrease by TCDD in Lgr5 mRNA levels in vitro. These results suggest that RSPOs bind to LGRs located in the BE of the UGS to initiate prostate bud formation. Together, these data illustrate that TCDD inhibits multiple components of the Wnt signaling pathway and that the combined inhibition of these components significantly contributes to the inhibitory budding phenotype caused by TCDD. (Grant support: NIH ES01332, T3Z ES007015)

2142 Sensitivity of Toxicological Endpoints to Detect Alterations in the Male Reproductive System of Nonhuman Primates.

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The assessment of the potential for a compound to cause toxicity to male reproductive processes relies heavily on histopathology and organ weights of the reproductive tissues. This is even more so with biologics where non-human primates (NHP) is the only relevant species and evaluation of functional fertility is impractical. To address this, this study used an ICH S6(R1) included a trigger for the addition of non-conventional endpoints to NHP studies in cases where there is reproductive cause for concern. To determine the sensitivity of these endpoints to perturbation we performed a power analysis of routine and triggered endpoints. The analysis was done on control data from sexual maturation Asian and Mauritian sourced NHP used in toxicity studies at Covance Laboratories GmbH. The power calculations were performed with a 2 sample two-sided T-test (a=0.05) assuming 3 NHP/group and the number of observations ranged from 98 to 472 per endpoint. For reproductive organ weights the power to detect a 50% change from control was 30%, 35%, 18% and 13% for testes, epididymides, prostate, and seminal vesicle, respectively. For testicular volume the power to detect a 50% change was ~30%, with slight differences depending on the method (caliper or ultrasound). For seminal analysis the power to detect a 50% change was 6%, 6%, 60% and 41% for ejaculate weight, sperm count, sperm motility, and sperm morphology (percent normal), respectively. For male hormone data the power to detect a 50% change was 18%, 30%, 7% and 78% for testosterone, inhibin B, luteinizing hormone, and follicle stimulating hormone, respectively. Given samples sizes of 3 per treatment group, and the magnitude of biological variability observed for these endpoints, the ability to draw conclusions about potential toxicity will be limited for most endpoints. Consequently, great care should be taken in the choice of endpoints added to general toxicity studies to assess the male reproductive system.

2143 Contribution of PI3K/Akt/DAF-16 Activity in C. elegans to Gene-Environment Interactions following MeHg Exposure.

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Neither genetic nor environmental factors completely explain the etiology and progression of dopaminergic (DAergic) neurodegeneration in Parkinson’s disease (PD). This study examines PI3K/Akt signaling as a mechanistic link between environmental and genetic factors known to contribute to PD. Specifically the interaction between methylmercury (MeHg) exposure, a ubiquitous neurotoxin, and PARK7 (DJ-1), an autosomal recessive PD gene. PI3K/Akt signaling is highly conserved and regulates many aspects of neuronal survival and function. MeHg exposure has been shown to activate PI3K/Akt signaling and DJ-1 modifies PTEN activity, an upstream regulator of PI3K/Akt. Mutant and transgenic Caenorhabditis elegans (C. elegans) strains were used to determine if PI3K/Akt responsiveness and loss of DJ-1 increases MeHg vulnerability. Strains used include; wild type (WT) Bristol N2; strains null for daf-16, age-1, daf-18, djr-1.1, and djr-1.2 and a strain overexpressing DAF16::GFP. Synchronous L1 populations were treated with 0-40μM MeHg for 30 minutes followed by assessment of survival and basal slowing (a dopamine-dependent feeding behavior). Compared to WT worms, increased PI3K/Akt activity resulting from genetic loss of daf-18 (PTEN homolog) anddaf-16 (FOXO homolog) led to increased death and lower basal lethal dose (DL50) values following MeHg treatment while decreased PI3K/Akt activity through loss of age-1 (PI3K) homolog and overexpression of DAF-16::GFP increased survival. Loss of djp-1 or djp-1 (PARK7 homolog) did not affect survival. Basal slowing was significantly reduced in WT worms following 20μM but not 10μM MeHg treatment (p<0.01). Slowing in worms lacking djp-1 but not djp-1,2, was decreased by 10μM and 20μM MeHg, suggesting that loss of djp-1 increases DAergic sensitivity to MeHg. Collectively, these findings demonstrate that MeHg-induced
Methylmercury (MeHg) hastens the onset of motor paralysis in SOD-1 G93A mice, a genetic model of Amyotrophic Lateral Sclerosis (ALS), suggesting a gene-environment interaction in a genetically susceptible animal model. Associated with this are alterations in [Ca2+]i, and glutamate receptor (GluR) function, both of which are suggested targets for MeHg toxicity. The goal of this study was to determine if MeHg alters mRNA levels of voltage gated Ca2+ channels (VGCCs), NMDA, or AMPA receptors. We measured mRNA levels of the pore forming α subunit of L, N, P/Q and R type VGCCs, NR1 and NR2A subunits of NMDA receptors and GluR2 and GluR3 subunits of AMPA receptors. Determining if MeHg alters levels of these ion channels could contribute to understanding of MeHg-induced alteration in [Ca2+]i, in motor neurons. Postnatal day 5 rats were treated sc with 0.75 or 1.5 mg/kg/day MeHg for 15 or 30 days (d), then stopped for a clearing period of 30 d. Quantitative real time PCR was performed on reverse transcript of RNA isolated from 10 mg of brainstem tissue, a region rich in motor neurons. MeHg treated animals had higher levels of GluR2, GluR3 and NR2A at 15 d in both 0.75 and 1.5 mg/kg/day exposure. MeHg induced an increase in the levels of all GluR subunits studied. It was highest at 30 d, and more pronounced in the 0.75 mg/kg/day exposure. GluR2 was the most affected subunit. At 60 d, levels of all GluR subunits were increased but at levels similar to those seen at 15 d of exposure. With the exception of α1, levels of all VGCC α subunits were higher at 0.75 mg/kg/day MeHg exposure on 15 and 30 d and lower at 60 d. At 15 d 1.5 mg/kg/day MeHg increased levels of all VGCC α subunits except α2C. At 30 d the level was even higher for all of the α subunits, but returned to levels similar to those at 15 d after the clearing period. The increase in the level of GluR subunits and most of the VGCCs could contribute to the alteration in Ca2+ homeostasis induced by MeHg on motor neurons. Supported by NIH grants R01ES03299 and R2NS0657777.

Methylmercury (MeHg) causes marked alterations in neuronal [Ca2+]i homeostasis through its interaction with membrane ion channels such as voltage-gated Ca2+ channels (VGCCs), and Ca2+ permeable ligand-gated-NMDA (NMDAR) and AMPA receptors (AMPAR). Chronic postnatal exposure to MeHg hastens the onset of Amyotrophic Lateral Sclerosis-like phenotype in the SOD1G93A-A genetically susceptible mouse model via glutamate-mediated excitotoxicity. The underlying mechanism of MeHg effects on brainstem motor neurons remains unclear. In this study we focused on examining the effects of chronic MeHg exposure on mRNA expression of the pore-forming α subunits of the L (α1), N (α2C), P/Q (α3), and R-type (α1D) VGCCs, NR1 and NR2B subunits of the NMDAR, GluR2 and GluR3 subunits of the AMPAR, Ca2+ binding proteins calbindin D28K and parvalbumin, and glutamate transporter EAAT2, all of which can modulate [Ca2+]i homeostasis. Eleven week old male ICR mice were exposed to 0, 0.5 or 5 ppm MeHg in drinking water ad libitum for 6 mos. Total RNA was extracted from 10 mg of brainstem and reverse transcription PCR performed. Quantitative real time PCR was performed on reverse transcript (cDNA) to measure expression of the target genes. Exposure to 0.5 ppm MeHg induced a decrease in the expression of the AMPAR subunits, Ca2+ binding proteins and the α1 subunit of the N- and P/Q-type VGCCs. NMDAR expression at 0.5 ppm MeHg exposure was not altered at the NR1 subunit but was increased for NR2B. At 5 ppm MeHg decreased expression of GluR2, EAAT2, calbindin D28K, parvalbumin, and all of the VGCC subunits. Expression of AMPAR subunit GluR3 was increased at 5ppm exposure. These results support the idea that MeHg-induced increase in [Ca2+]i on brainstem motor neurons is due to the glutamate receptors (greater GluR2 expression, and NR2B expression), or inefficient Ca2+ buffering (decrease in the Ca2+ binding proteins and EAAT2 transporter). VGCC may not contribute to the alteration in [Ca2+]i induced by MeHg because their expression was decreased. Supported by NIH grants R01ES03299 and R2NS0657777.

Methylmercury (MeHg) is an environmental pollutant that is toxic to the developing central nervous system (CNS) in children, even at low exposure levels. Perinatal exposure to MeHg is known to induce neurological symptoms with neuropathological changes in the CNS. However, the relationship between the neurological symptoms and neuropathological changes induced in offspring as a result of exposure to low-dose MeHg is not well defined. In the present study, neurobehavioral analyses revealed that exposure to a low level of MeHg (5 ppm in drinking water) during developmental caused a significant deficit in the motor coordination of rats in the rotating rod test. In contrast, general neuropathological findings, including neuronal cell death and the subsequent nerve inflammation, were not observed in the region of the cerebellum responsible for regulating motor coordination. Surprisingly, the expression of synaptophysin (SPP), a marker protein for synaptic formation, significantly decreased in cerebellar granule cells. These results showed that perinatal exposure to low-dose MeHg causes neurobehavioral impairment without general neuropathological changes in rats. We demonstrated for the first time that exposure to low-dose MeHg during development induces the dysfunction of motor coordination due to changes of synaptic homeostasis in cerebellar granule cells.

Methylmercury (MeHg) is a potent environmental neurotoxicant that causes cell-type specific damage in the cerebellum in a calcium (Ca2+) dependent manner. The developmental effects of MeHg have been widely explored. However, the effects of chronic exposure in adults have not been characterized. The ability of MeHg to cause cell-type specific damage due to alterations in Ca2+ regulation raises the possibility that other subsets of neurons with vulnerability that depends on Ca2+ regulation could be sensitive to MeHg. This study focused on investigating the effects of chronic MeHg treatment on the mitochondrial function of nigrostriatal dopamine (NSDA) neurons. These neurons exhibit a unique physiological phenotype; they autonomously generate action potentials in the absence of synaptic input. The spontaneous action potentials rely on the influx of Ca2+ through Cav1.3, L-type Ca2+ channels. We also investigated whether co-treatment with isradipine, a Ca2+ channel antagonist, would protect against MeHg-mediated damage. Beginning at 3mo, male BALB/c mice were given 5ppm Hg as MeHg in their drinking water alone or co-treated with 2ppm isradipine in their food for 6mo. Following the 6mo treatment period mouse weight gain was normal and there were no phenotypic changes. Mitochondrial function was measured in synaptosome preparations from NSDA neurons using the Extracellular Flux Analyzer (Seahorse Biosciences). Mitochondrial basal respiration, ATP production rate, maximal respiration, and spare capacity were examined. No alterations were seen in any measure of mitochondrial function in animals treated with MeHg alone or in animals co-treated with isradipine (n=9, p-values=0.8494, 0.1019, 0.9178, 0.8227 respectively). These studies have demonstrated that mitochondrial function of striatal synaptosomes is unchanged following a 6mo MeHg treatment. The effects of a 12mo MeHg treatment are currently being pursued. This work was supported by a VICTER supplement to R01ES03299.

Methylmercury (MeHg) exposure from occupational, environmental, and food sources is a significant threat to public health. Recent epidemiological and vertebrate studies suggest that MeHg exposure may contribute to the propensity to develop Parkinson’s disease (PD). We have developed a novel C. elegans model of MeHg toxicity and have shown that low, chronic exposure confers dopamine (DA) neuron degeneration, and that the toxicity is partially dependent on the phase II antioxidant transcription factor SKN-1/Nrf2. Aims/Objectives: In this study we
asked what genes are involved in MeHg-induced animal and DA neuron pathology. Methods: We utilized a reverse genetic screen that has identified a number of molecular transporters and proteins involved in MeHg resistance. Support: NIEHS.

2149 Biochemical Alterations of Rat Brain Mitochondrial Enzymes Induced by Aluminium Chloride.
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Aim: The present study was planned to investigate the effect of aluminium chloride (AlCl3) on enzyme activities in the mitochondrial fractions of the brain of male albino rats. Methods: Adult male albino rats were administered AlCl3 at two different doses, 50 mg and 100 mg/kg body weight, orally, daily for 45 days. At the end of the experimental period the animals were sacrificed, the brain was removed and mitochondrial fractions were isolated. Antioxidant enzymes like catalase, superoxide dismutase, glutathione peroxidase, glutathione reductase and glutathione S-transferase were estimated in the brain extract. Other biochemical markers were also studied. Results: AlCl3 administration had no effect on the weight and brain weight. Almost all the antioxidant enzymes studied were markedly diminished in the brain of AlCl3 treated animals. The activities of acid phosphatase and alkaline phosphatase were significantly increased. The activities of Ca++ ATPase, Mg++ ATPase and Na+ K+ ATPases were decreased by the AlCl3 treatment in the brain. However, the influence was found to be more in 100 mg treated when compared to 50 mg AlCl3 treated rats. The activity of acetylcholinesterase was diminished while the lactate dehydrogenase activity increased after aluminium treatment. Conclusion: The present study suggests the toxicity of aluminium by inducing the oxidative stress and adverse alterations in the brain metabolism and possible interference in brain coordination processes.

2150 Effects of Mercuric Chloride on Cell Surface Expression of Dopamine Transporter in PC12 Cells.

Mercury compounds are known to have a ubiquitous nature due to their ability to move through the biogeochemical cycle and food web. Organic and inorganic forms of mercury generally target different organ systems of the body while bio- transformation after exposure can result in the bioaccumulation of various species of the element. Divalent mercury (Hg2+) can be found within the brain following exposure to the mercury vapor of dental amalgams and after extended exposure to methyl mercury. Dopamine transporter protein (DAT) has been implicated as a link between neurotoxicity following exposure to 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP). Increases in DAT expression on the cell membrane have been shown to result in amplified sensitivity to MPTP. The in vitro effect of mercuric chloride (HgCl2) on the cell surface expression of DAT in stably trans- fected DAT expressing PC12 cells was studied. Cell viability (MTT assay) following treatment with 0.5ppm HgCl2 and 1-methyl-4-phenylpyridinium (MPP+) shows the active metabolite of MPTP was observed to result in an additive increase in toxicity compared with treatment with MPP+ alone. Using immunocytochemistry and Western blot, DAT surface expression was seen to increase following HgCl2 (0.2-2.0ppm) exposure which exhibited a biphasic pattern with the maximum expression observed at 0.5ppm when compared to the control. With null function DAT mutant, cell viability following exposure to HgCl2~2 and MPP+ showed an absence of the increase in toxicity observed at 0.5ppm originally observed in the wild type control cells. The results show that HgCl2~2 is able to increase DAT surface expression in a concentration dependent manner and as a result, exposure could trigger subsequent increased susceptibility to pesticides such as MPTP that use the DAT system as pathway of toxicity.

2151 Changes in Gene Induction Associated with Lead Acetate Mediated Oxidative Stress and Mitochondrial Dysfunction in Neuronal PC12 Cells.
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Lead is a major environmental heavy metal toxic that affects brain development and impairs cognition in children and adults. Despite extensive research, molecular mechanisms underlying neurotoxicity mediated by lead are not well understood. Previous studies have shown that ROS production caused by lead exposure decreases cellular antioxidant defense networks in the brain. In the present studies, we evaluated lead-mediated alterations in oxidative stress pathways in PC12 cells. Using H2DCFDA in conjunction with flow cytometry, we found that lead exposure increased ROS production up to 2-7 fold (3 h, 1 nM to 10 μM) of this effect was associated with mitochondrial membrane potential changes which were 60% lower after 24 hour treatment with 1 μM PbAc2. Using QPCR, we observed that exposure to lead resulted in the induction of anti-oxidant genes including MnSOD, catalase, NQO1, HO-1 and GST-M1. QPCR analysis also revealed induction of genes that are indicators of Ca2+ flux (TRPV1), inflammation (COX-2, IL-1 β, TNFα), and apoptosis (bcl-2, bax, caspase-3). Relative mRNA for signaling kinases ERK1/2 (p44), SAPK/JNK, and p38 were increased while that for p38MAPK was decreased. Taken together our results suggest that lead treatment of PC12 cells enhances cellular ROS levels, negatively impacts the anti-oxidant gene profile and the mitochondrial membrane potential of PC12 cells, upregulates expression of genes critical for inflammation and apoptosis and alters expression of critical cellular signaling kinases. We speculate that changes in the expression of antioxidant enzymes and signaling kinases are important in lead-mediated neurotoxicity. (Supported by AR055073)

2152 Effects of Pb2+ in the Neural Differentiation of Mouse Embryonic Stem Cells.
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Exposure to environmental agents during embryonic life is suspected to modify the epigenetic mechanisms that regulate the gene expression patterns that control development. The resulting changes in gene expression may affect lineage differentiation and extend into adulthood, well beyond the time the organism was exposed to the agents. This paradigm provides a testable molecular basis for the Barker hypothesis, which proposes that there is a fetal origin of adult disease. Lead (Pb) is a ubiquitous environmental toxicant whose possible effects, especially at early ages of development, include reduction of cognitive functions and IQ, behavioral effects, attention deficit and hyperactivity disorder. We are using neural differentiation of mouse embryonic stem cell (mESC) to determine the molecular changes taking place during embryogenesis and neurodevelopment as a consequence of long-term exposure to Pb2+.

We find that after differentiation of mESC into neuronal cells, cultures express many neuronal markers, including Tubb3, Syp, Gap43 and Hud, and do not express the glial marker Galc. Furthermore, the cells express Vglut1, a marker of glutamatergic neurons. Incubation with Pb2+ during differentiation reduces the expression of the calcium-dependent exon IV of Bdnf and alters the expression pattern of other neuronal gene but does not alter the morphology of the neuron and the expression of Tubb3. Thus, we conclude that after differentiation of mESC in vitro, we can obtain pure cultures of glutamatergic neurons whose gene expression patterns are changed by Pb2+ exposure. These cells should provide an useful substrate to analyze the mechanisms of toxicity of environmental neurotoxins. Supported by NIH grant R21 ES020948.
Developmental Lead (Pb2+) Exposure and Mutant DISC1 Interact to Produce Schizophrenia-Like Neurobehavioral Abnormalities and Brain Volume Changes in Mice: A Gene-Environment Interaction Study.


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The glutamatergic hypothesis of schizophrenia suggests that hypoactivity of the NMDA receptor (NMDAR) is an important factor in the pathophysiology of schizophrenia and related mental disorders. The environmental developmental neurotoxicant lead (Pb2+) is a potent and selective antagonist of the NMDAR and two recent human studies have suggested an association between prenatal Pb2+ exposure and the increased likelihood of expressing a schizophrenic phenotype later in life. Schizophrenia and other major mental disorders likely result from interactions between genetic risk factors and adverse environmental insults. We evaluated the neurobehavioral consequences of Pb2+ exposure in mice with inductive expression of mutant Disrupted-in-Schizophrenia-1 (DISC1), a candidate gene for major psychiatric diseases. We hypothesize that mutant DISC1 and Pb2+ exposure synergistically interact to produce an exaggerated phenotype in mutant mice. Mutant DISC1 and control mice born by the same dams were raised and maintained on normal or Pb2+ diet. We tested animals in a series of behaviors associated with schizophrenia and performed volumetric MRI measurements of the brain. We found that the interaction between developmental Pb2+ exposure and mutant DISC1 produced a significantly greater increase in locomotor activity, impaired pre-pulse inhibition of the acoustic startle that was abrogated by D-serine and exacerbated responses to MK-801, an NMDAR antagonist. Compared to control unexposed mice, mutant exposed animals had significantly larger lateral ventricles. Our data suggest that interactions between Pb2+ and mutant DISC1 may synergistically produce schizophrenia-like behavioral alterations and brain volume changes in mice. [Supported by grant # ES06189-18S1 to TRG]

Reduced Parvalbumin Expression in the Striatum, Frontal Cortex, and Hippocampus after Developmental Pb2+ Exposure: Examining Early Life Pb2+ Exposure As a Risk Factor for Schizophrenia.

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The calcium-binding protein, parvalbumin (PV), is expressed in GABAergic interneurons and is implicated in working memory and associative learning. Decreased PV expression is observed in schizophrenia patients and also in animal models of schizophrenia. N-methyl-D-aspartate receptor (NMDAR) antagonists are used to model certain aspects of schizophrenia in animal models and NMDAR antagonists decrease PV levels and the number of PV-positive interneurons, suggesting that NMDAR function plays a role in the expression of PV interneurons and in schizophrenia. Lead (Pb2+) is a potent NMDAR antagonist that has been implicated in the etiology of schizophrenia, therefore we examined the effect of developmental Pb2+ exposure on PV cell density in the striatum and PV protein expression in the striatum, hippocampus and frontal cortex of rats developmentally exposed to Pb2+. A trend of decreased PV positive cell density was observed at all levels of the striatum, however a statistically significant decrease (t(11)=2.2, p = 0.005) in cell density was found in the caudal striatum of Pb2+ treated rats relative to controls. Additionally, PV protein expression measured by western blot was significantly reduced in the striatum (t(22)=5.3; p = 0.003), frontal cortex (t(7)=2.25, p = 0.05) and hippocampus (t(10)=2.31, p = 0.04) of rats developmentally exposed to Pb2+. Overall, these findings indicate that early life Pb2+ exposure decreases the number of PV positive neurons in the striatum, and reduces the expression of PV in the striatum, frontal cortex and hippocampus. The data suggests a relationship between Pb2+ induced NMDAR hypoactivity and PV expression, implicating a potential association between developmental Pb2+ exposure and the expression of a schizophrenia phenotype later in life. [Supported by NIEHS grant # ES06189 to TRG]

Effects of Developmental Lead Exposure on Associative Learning and Memory Are Modified by Sex, Developmental Window of Exposure, and Level of Exposure.

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Trace fear conditioning is a variant of fear conditioning in which a neutral conditioned stimulus (CS) is paired with an aversive unconditioned stimulus (UCS). Trace conditioning includes a “trace” interval of several seconds separating the CS and UCS. Learning the CS–UCS association across this interval and the consolidation of the memory requires participation of both the hippocampus and medial prefrontal cortex (mPFC). Long-term storage of information associating temporally disconnected events occurs in the mPFC in parallel with memory storage in the hippocampus. The current research investigated the extent to which these processes are affected by developmental Pb exposure. Long Evans dams were fed Pb-containing food (RMH 1000 with or without added Pb acetate: 0, 150, 375, 750 ppm) prior to breeding and stayed on the same diet through weaning at postnatal day 25 (perinatal exposure group (Peri)). Other animals were exposed to the same doses of lead but exposure started on postnatal day 1 and continued through postnatal day 25 (early postnatal exposure group (EPN)). Beginning at postnatal day 45, animals were placed in the trace fear conditioning apparatus. Conditioning trials (CS tone–UCS shock pairings) were repeated six times during an 18 min training period. Freezing behavior was measured during the trace period. Retention testing occurred 24/48 hrs and 10 days later to assess memory consolidation and long-term memory. At the lowest level of exposure, EPN-exposed females were more impaired at short and long-term retention than were Peri-exposed females; the opposite effect was observed in males. In females, the lowest level of exposure had the greatest disruptive effect on retention. In males, the highest level of exposure had the greatest disruptive effects on retention. These data suggest complex responses of the brain to developmental Pb exposure with likely different molecular effects in hippocampus and mPFC that vary with sex and timing and level of exposure. Supported by NIH ROI-E5015295.

Sex- and Hemisphere-Dependent Neurochemical Changes Produced by Lead, Prenatal Stress, and the Combination.

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Brain lateralization, important to mediation of cognition and ‘multi-tasking’, is disrupted in attention deficit disorder and schizophrenia. Altered brain lateralization could play a role in the corresponding cognitive and attention deficits associated with both low level lead (Pb) exposure and prenatal stress (PS). This study examined laterality of mesocorticolimbic (frontal cortex, nucleus accumbens, striatum, midbrain) monoamines and amino acids (frontal cortex only) that mediate such behavioral functions, both under normal (control) conditions, and in response to lifetime Pb (0 or 50 ppm), PS (restraint stress on gestational days 16-17) or Pb+PS in rats. Sex-dependent differences in brain laterality were seen even in control rats, as noted in frontal cortex, striatum and midbrain monoamines and frontal cortex amino acids. Hemispheric differences in Pb ± PS effects were most notable in males, as seen in frontal cortex and striatum, but particularly in midbrain, where Pb+PS, but neither alone, significantly reduced right hemispheric levels of homovanillic acid and norepinephrine, with similar trends in serotonin. In contrast, no Pb+PS effects were found in left hemisphere. Pb+PS, but neither alone, reduced frontal cortex dopamine and increased dopamine turnover in left hemisphere of males, whereas in the right hemisphere, Pb, Ps and Pb+Ps reduced frontal cortex dopamine and Pb alone increased dopamine turnover. The only suggestive Pb+PS hemispheric difference in females was increased right but not left hemisphere frontal cortex GABA. Thus Pb, Ps and Pb+Ps can differentially influence mesocorticolimbic neurotransmitter function in female rats, and Pb+PS effects are likely to contribute to associated sex-related alterations in behavioral outcomes, such as our previously reported contrasting effects on learning accuracy produced by Pb+P in males vs. females. The findings also underscore the significance of defining the hemisphere used for assay in experimental studies for evaluation of CNS mechanisms of behavior.

Sex-Dependent Changes in the Effects of Lead and Prenatal Stress on Impulsivity and Neurochemical Substrate.

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Both lead (Pb) exposure and prenatal stress (PS) adversely affect cognition and attention... Impulsivity is one diagnostic component of attention deficit, and enhanced impulsivity has been related to multiple neuropsychiatric disorders. This
study examined whether combined Pb+PS would enhance impulsivity compared to either alone, as PLE can alter surface area and volume (abundance) decreased in spherules and increased in pedicles. PLE remodeling of spherule mt produced smaller cristae with more branching, whereas pedicle mt had larger cristae with more branching and increased cristae junction diameter. PLE decreased dark-adapted PR and PR synaptic terminal QO2. In Bcl-xL/PLE, spherules still had decreased QO2 while pedicles had increased branching, cristae segments/volume, and cristae junction diameter; and PR and synaptic QO2 only partially recovered.

These findings reveal cellular and compartmental differences in the structure, vulnerability and remodeling of rod and cone inner segment and synaptic mt to PLE: consistent with findings that synaptic mt are more sensitive to calcium overload, oxidative stress and ATP loss than non-synaptic mt. These PLE alterations likely underlie the persistent scotopic-mesopic deficits, and stress the importance of examining synaptic dysfunction after developmental insults and preventing synaptic degeneration even if apoptosis is blocked.

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2160 Alpha-Synuclein-Mediated Activation of c-Abl and Dopaamine Depletion in Dopaminergic Neuronal Cells Treated with Iron-Oxide Nanoparticles or Methamphetamine.

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Mutations in the alpha-synuclein gene have been associated with autosomal dominant forms of Parkinson’s disease (PD). Transgenic mice that over-express the human alpha-synuclein gene (primarily the point mutations A53T and A30P), develop neurological impairments similar to those of PD. Previous studies in our laboratory have shown that oxidative-stress-mediated activation of the tyrosine kinase, c-Abl, results in an increase in the phosphorylation of parkin, an important E3 ubiquitin ligase that assists in the clearance of proteins destined for proteasomal degradation. Here, we show that treatment with iron-oxide nanoparticles or methamphetamine results in activation of c-Abl, observed via the measurement of phospho-Abl. Additionally, an over-expression of alpha-synuclein protein in SHSY5Y neuroblastoma cells was observed after these treatments. A 45% increase in the expression of alpha-synuclein was observed in SHSY5Y cells treated with iron oxide nanoparticles (10 and 30 nanometers) at a concentration of 10 μg/ml. Similarly, a 55% increase in the expression of alpha-synuclein protein was observed 24 h after exposure to 500 μM methamphetamine. In addition, a significant depletion in dopamine was observed after treatment with iron oxide nanoparticles or methamphetamine (55% and 65%, respectively), suggesting that both the over-expression of alpha-synuclein and exposure to oxidative stress, which is one of the major pathways of c-Abl activation. These results suggest that the over-expression of alpha-synuclein and dopamine depletion after exposure to iron oxide nanoparticles or methamphetamine contribute to oxidative-stress mediated activation of c-Abl, thus initiating events that may lead to dopaminergic neuronal cell death.


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Alzheimer disease (AD) is the most common form of dementia affecting around 24.3 million people worldwide. As a consequence of the rapid demographic ageing, AD has become one of the most severe progressive socio-economic and medical burdens facing countries all around the world. AD brains are characterized by the presence of extracellular deposits of amyloid-ß-containing plaques and intracellular neurofibrillar tangles (NFTs) composed of paired helical filaments of hyperphosphorylated Tau. AD is considered to be the result of complex events involving both genetic and environmental factors. Among them, two remarkable factors are oxidative stress and mitochondrial damage. Postmortem examinations of human brain tissue have demonstrated that iron concentrations are increased in the brains of AD patients compared to controls. The main goal of this study was the identification of proteins showing different expression levels in AD, using iTRAQ-protein labeling.
Drug induced liver injury (DILI) is a leading cause for drug failures in clinical trials and for post-approval drug withdrawals. This fact implies that the current animal-based approaches do not provide a sufficiently reliable assessment of DILI; therefore, there is an increased interest to develop new approaches for identifying hepatotoxicity, especially those that can be applied in the early drug discovery. In this study, we assessed the feasibility of using high content screening (HCS) assays of rat primary hepatocytes to predict the human specific hepatotoxicity of pharmaceuticals. We developed a predictive model using a commercial HCS assay on 8 cellular parameters that were relevant to hepatotoxicity mechanisms including cell loss, nuclear size, DNA damage, apoptosis, lysosomal mass, DNA fragmentation, mitochondrial potential, and steatosis. The model’s performance was assessed both by internal cross-validations using a library of 108 drugs with human-specific hepatotoxicity annotations and by further external validation using an additional sixteen drugs. In the leave-one-drug-out cross-validation, the HCS assay-based predictive model yielded accuracy of 68% with sensitivity of 77% and specificity of 50%, which is significantly higher than that by chance and better than the quantitative structure–activity relationship (QSAR) model developed using the same set of drugs in accuracy. The results from the external validation showed that eight out of nine of the most-DILI-concern drugs (with 89% sensitivity) and four out of seven DILI-concern drugs (with 57% specificity) were correctly predicted. Our findings suggest that the HCS assay of rat hepatocytes can potentially approach useful toxicology needs to be further elucidated.

Scope: The amount of preclinical research based on peptides is constantly growing, however the difficulties of reliably analyzing this type of molecule leads to a major challenge in obtaining reliable toxicokinetic data. Combining the latest technology and, identification and quantitation of peptides by tandem mass spectrometry. We found up-regulated such as Ferritin light chain, Ferritin heavy chain and Melatonin. In addition, several proteins involved in redox regulation also identified more than 721 polypeptides, where some iron regulatory proteins were gene product and downregulated. The relationship between physicochemical properties and compound accumulation was derived by the low pH in the lysosomes. In addition our data revealed membrane perturbation by the compounds with similar physicochemical properties. In an effort to understand how physicochemical properties can contribute to mitochondrial compound accumulation we selected a set of basic compounds and measured their logD (partition coefficient) at neutral (7.4) and acidic (4.7) pH. The majority of basic compounds with acidic logD (at pH 4.7) > 0.3 induced more toxicity in HepG2 cells cultured in galactose than in cells cultured in high glucose-containing media, indicating mitochondrial liability associated with these compounds. This data supports the hypothesis that compounds with relatively pH-insensitive permeability selectively accumulate into mitochondria. Interestingly, certain compounds such as nefazodone and nardilipine were shown to carry both, lysosomal and mitochondria liabilities. In summary, our data indicate that physicochemical properties alone can contribute to both, lysosomal and mitochondrial compound accumulation. How this contributes to in vivo organ toxicity needs to be further elucidated.

The β-site amyloid precursor protein (APP)-cleaving enzyme 1 (BACE1) is a target of tremendous focus and development interest for the treatment of Alzheimer’s disease. While initial characterization reports of BACE1 knockout (KO) mice suggest that BACE1 is a safe target for modulation (Roberts et al., 2001; Lou et al., 2001), more recent studies present evidence of retinal pathology (Cai, et al., 2012). Additionally, oral administration of a selective BACE1 inhibitor to rats has been associated with retinal changes after chronic dosing, including accumulation of autofluorescent material in the retinal pigment epithelium (RPE) and degeneration of photoreceptors in associated retina (May, et al., 2011). To address these concerns we evaluated the retinas in BACE1 homozygous KO mice from an in-house colony. With the goal of enabling a shorter duration screening paradigm for ocular toxicity, we also evaluated the retinas from rats after a single intravitreal injection of BACE1 inhibitors at non-specific protease inhibitor at high (-10 μM) intracellular concentrations. We found that 3-month old BACE1 KO mice are viable and without organ or clinical pathology distinctions from wild-type (wt) littermates. Ocular evaluation by H&E staining indicated that the retinas of the BACE1 KOs were unremarkable when compared to wt mice. Ophthalmic examinations, as well as light and fluorescent microscopy following intravitreal injections of BACE1 inhibitors, revealed no significant differences in the retinas or RPE between control and treated rats 1-8 days post-injection. In summary, our results support those of others indicating retinal pathologies reported with BACE1 inhibitors are not related to downregulation of BACE1 activity. Additionally, short duration intravitreal exposure to high concentrations of selective BACE1 inhibitors or non-specific protease inhibitor was not associated with retinal pathology. This finding may not be a reliable model for predicting retinal toxicity of BACE1 inhibitors.
related to dried blood spots and mass spectrometry provides many advantages even though the initial method development is more arduous than for typical small molecules of less than 1000 daltons.

Experimental Procedures: The peptide calcitonin-salmon was chosen based on it being readily available from Sigma Aldrich and that its molecular weight is higher than normally monitored by mass spectrometry since it consists of thirty-two amino acid linear polypeptide. The compound is dosed by oral gavage at 10 mg/kg to CD-1 mice, and 8 ml of blood collected by tail clipping every two minutes. The blood is immediately transferred to Whatman 903 filter paper and allowed to dry before placing with desiccant and refrigerating. The dry blood samples are then punched out, and calcitonin-salmon is extracted from the spots and injected. The chromatographic conditions were developed using reverse phase chromatography and detection using a triple quadrupole API 4000 mass spectrometer with electrospray ionization to monitor the parent and fragment ions of both calcitonin-salmon and its analog internal standard.

Conclusions: Results show that this approach provides a toxicokinetic profile with timepoints every two minutes whilst allowing a considerable reduction in mice required for a study and giving a full profile for each individual animal. This work shows that preliminary TK data may be obtained based on 3R principles even for peptide analysis, an area that still has room for much improvement using dry blood spot sample analysis.

2167 Early and Progressive Changes in the Urinary Biomarker Profile of Cisplatin-Induced Kidney Injury in Cynomolgus Macaques.

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Cisplatin-induced nephrotoxicity has been characterized in animals and humans. New urinary biomarkers may enable early or parallel detection of drug-induced kidney injury (DIKI) in non-rodent models. Our aims were to: 1) determine if a human renal biomarker panel can detect early DIKI when compared with traditional biochemical and histological indices; 2) monitor progressive cisplatin-mediated changes in DIKI biomarkers over 20 days in cynomolgus macaques. Animals (3M/3F per group) were treated with a single dose of cisplatin (2.5 mg/kg i.v. infusion; groups 1/2/4/5) or saline (groups 3/6) on day 1. Toxicokinetic profile was determined on day 1. Blood and urine samples were collected predose and postdose (days 1, 4, 9, 15, 20) for comparing changes in plasma chemistry, urinalysis and urinary DIKI biomarkers. Renal tissue was examined microscopically at different intervals. Results indicated cisplatin produced progressive renal histologic changes from sublethal findings on day 2 to maximum effects on day 21. Traditional renal markers significantly increased in plasma creatinine (8-fold), BUN (3-fold), urinary glucose (5-fold), protein, and enzymes (GGT, NAG) day between 4-9 but resolved by day 15 when renal pathology appeared more severe. Four new biomarkers (clusterin, calbindin, lipocalin, Tamm-Horsfall protein) changed progressively and significantly from 4 to 20 days to parallel development of renal histologic injury. Importantly, β2-microglobulin was significantly reduced before substantial renal damage observed on day 2 and increased progressively over 20 days to parallel histologic changes, suggesting this is the most sensitive biomarker for detection of onset and potentially prodrmal cisplatin-induced kidney injury in cynomolgus macaques.

2168 A Case Study for Exploring Rodent Models to Assess Risk of Hypotension.


A cardiovascular (CV) safety pharmacology study in telemetered conscious dogs evaluating Compound-A (CPD-A), an anti-cancer agent, confirmed a dose- and time-dependent decrease in mean arterial blood pressure (MAP), most likely attributable to off-target PDE3 inhibition. At 20 mg/kg p.o. CPD-A, MAP was reduced by ~30% from the baseline within 1-hour of dosing and this effect persisted ~12 hours. The objective of this study was to identify a rodent model that would best mimic the dog CV response to CPD-A. We investigated CPD-A (at doses up to 120 mg/kg, p.o.) in conscious, telemetered rats and detected no meaningful acute effects on MAP although a moderate tachycardia was observed. Upon repeated daily dosing of 120 mg/kg QD to rats, CPD-A led to a mild decrease in MAP on Day 3 when compared to pre-dose baseline and time-matched vehicle controls. As in the acute study, a moderate tachycardia was observed. To maximize exposure, a single dose of CPD-A (30 mg/kg, p.o.) was administered in conscious rats and CPD-A was investigated. This resulted in a significant MAP drop and a moderate reflex tachycardia. Based on this data, we hypothesized that a difference in sympathetic tone and baroreflex sensitivity between dog and rat may underlie the species difference in cardiovascular response observed after single oral administration of CPD-A. To explore this possibility, we also assessed CPD-A in isoflurane-anesthetized rat model to decrease sympathetic tone and baroreflex response. Continuous infusion of CPD-A at 10 and 20 mg/kg/hr led to a gradual decrease in MAP of up to 15 and 45%, respectively, and a mild bradycardia at the highest dose. As an alternative to the rat, we also evaluated a urethane-anesthetized guinea pig model. Infusion of 10 mg/kg CPD-A over 30 min resulted in a gradual drop in MAP accompanied by reflex tachycardia. These studies suggest the anesthetized rat or guinea pig offered good in vivo screening alternatives for this compound class.

2169 MDR1, MR2 and BCRP Transporter Gene Knockout in CACO-2 Cells.

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Sponsor: Sigma-Aldrich, St. Louis, MO.

Purpose: Membrane transporters P-glycoprotein (P-gp, MDR1), MR2, and BCRP play a role in drug disposition and can mediate drug-drug interactions leading to safety/toxicity concerns. The goal of this study was to selectively knock out these drug transporter genes in a subclone of Caco-2 cells (C2BB1e) using zinc finger nuclease (ZFN) technology.

Methods: ZFN pairs targeting specific transporter genes were nucleofected into C2BB1e cells. Stable clones were isolated and sequenced at gene target sites. Transport function in wild type (WT) and knockout (KO) cell lines was tested in bidirectional assays using probe substrates at 5 μM for 2 hr at 37°C. Contents of both wells were analyzed for drug content using LC/MS-MS, and permeability was calculated for both directions. Lucifer yellow permeability was measured to verify monolayer integrity.

Results: ZFN-transfected clones were sequenced and the genotype was confirmed for disruptions in each allele. In bidirectional transport assays to confirm loss of transporter function, the efflux ratios (ER) for digoxin and erythromycin were reduced from 17.7 and 16.8 in the WT cells to 1.4 and 1.0 in the MDR1 KO cells, respectively. The ER for estrone 3-sulfate and nitrofurantoin were reduced from 22.7 and 13.2 in the WT to 1.8 and 1.7 in the BCRP KO cells, respectively. The ER for 5(6)-carboxy-2,7-dichlorofluorescein was reduced to 2.0 in the MR2 KO from 32.0 in the WT cells. Double knockout cell lines (MDR1/BCRP KO, MDR1/MRP2 KO, and MRP2/BCRP KO) were used to identify substrates of multiple transporters. Cimitidine was confirmed as a substrate of both P-gp and BCRP using the MDR1/BCRP KO cell line. Fexofenadine transport required P-gp and a basolateral MRP transporter such as MRP3, but not MR2.

Conclusions: Stable single and double transporter KO cell lines in the acute study, a moderate tachycardia was observed. To maximize exposure, a single dose of CPD-A (30 mg/kg, p.o.) was administered in conscious rats and CPD-A was investigated. This resulted in a significant MAP drop and a moderate reflex tachycardia. Based on this data, we hypothesized that a difference in sympathetic tone and baroreflex sensitivity between dog and rat may underlie the species difference in cardiovascular response observed after single oral administration of CPD-A. To explore this possibility, we also assessed CPD-A in isoflurane-anesthetized rat model to decrease sympathetic tone and baroreflex response. Continuous infusion of CPD-A at 10 and 20 mg/kg/hr led to a gradual decrease in MAP of up to 15 and 45%, respectively, and a mild bradycardia at the highest dose. As an alternative to the rat, we also evaluated a urethane-anesthetized guinea pig model. Infusion of 10 mg/kg CPD-A over 30 min resulted in a gradual drop in MAP accompanied by reflex tachycardia. These studies suggest the anesthetized rat or guinea pig offered good in vivo screening alternatives for this compound class.

2170 Target Promiscuity and Physicochemical Properties Contribute to Pharmacologically-Induced ER-Stress.

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In vivo toxicity of drug candidates remains a major problem in the pharmaceutical industry, and is a significant cause of late stage attrition. As a consequence predictive in vitro assays are developed and put in place early in the discovery pipeline to aid compound selection. Endoplasmic reticulum stress (ER-stress) has been implicated in many disease states, as well as compound-induced organ toxicities. We explored the role of ER-stress as a general mechanism of toxicity by utilizing a high-throughput in vitro assay to screen 318 chemically diverse Pfizer proprietary compounds with known in vivo toxicity outcome for nuclear accumulation of spliced X-Box Binding Protein 1 (XBP1s), a key transcription factor of the Unfolded Protein Response (UPR). We examined the correlation between physicochemical properties, such as molecular weight, pKa, lipophilicity, topological polar surface area, and passive permeability, as well as target promiscuity, between XBP1s hits and non-hits and found that lipophilicity, target promiscuity and low passive permeability significantly contributed to ER-stress. In addition, we have shown that compounds which cause ER-stress in the form of XBP1s activation at concentrations below 40μM have a more than four times greater chance of causing in vivo toxicity at 10μM plasma exposure.
2171 Enhancing the Safety of Antiviral Compounds by Assessing Mitochondrial and Nuclear Transcriptional Regulation Using a Multiplex Branched DNA Screen.

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The development of antiviral therapeutics can be hindered by the fact that they can cause damage to the liver, renal, and cardiac tissue. One antiviral strategy focuses on inhibiting the transcription of viral nucleic acids via nucleoside analog reverse-transcriptase inhibitors (NRTIs). NRTIs may cause toxicity due to mitochondrial DNA incorporation (mtDNA) and/or inhibition of DNA polymerase γ (Nature Rev Drug Disc 2003, 2: 812-22; Expert Opin Drug Metab Toxicol 2010, 6:1493-504). Functionally, such types of therapeutics may affect mtDNA transcription and replication, resulting in downstream defects in oxidative phosphorylation and mitochondrial biogenesis, which can lead to compromised organ function.

The strategies for ensuring mitochondrial safety include evaluating changes in mitochondrial biogenesis, metabolism, toxicity, and assaying for specific enzymatic activities in the electron transport chain. While these assays answer important functional questions, they may not necessarily pick up subtle changes with respect to DNA transcription and replication, which can lead to the downstream functional changes. The development of a multiplex, branched DNA (bDNA) assay to assess gene transcription in the mitochondria, as well as in the nucleus, has the advantage of detecting a total of eight transcripts in a single lysate. This significantly reduces the cost, time, and consumables that would be needed for traditional qRT-PCR. Studies using human liver epithelial cells treated with a selected group of nucleoside reverse transcriptase inhibitors, including 2,3'-dideoxycytidine, 2,3'-dideoxy-3'-thiacytidine, and 3'-azido-3'-deoxythymidine indicate that this technology may be applied to support the selection of compounds in pharmacological programs interested in pursuing nucleoside reverse transcriptase inhibitors as a part of their antiviral strategy.

2172 Comparison of In Vitro Models for Prediction of Hepatotoxicity of Pharmaceutical Drug Candidates.

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Liver toxicity is one of the foremost reasons for failure of a drug candidate to reach the clinic. It is also a leading cause for market withdrawals, FDA warnings and modifications of use for current medications. As a result, in vitro hepatocyte-specific studies using cells from preclinical species and also human donors are being investigated to allow for prediction of DILI during the early phase of drug discovery. During this project several novel in vitro models were studied for their potential to detect liver toxic compounds. A high-throughput, 384-well based assay with two-dimensional monocultures of rat or human primary hepatocytes was validated and used to detect liverotoxic compounds. A high-throughput, 384-well based assay with two-dimensional monocultures of rat or human primary hepatocytes was validated and implemented with ATP depletion as a cellular endpoint. Data obtained from moredimensional monocultures of rat or human primary hepatocytes was validated and used to detect liverotoxic compounds. A high-throughput, 384-well based assay with two-dimensional monocultures of rat or human primary hepatocytes was validated and used to detect liverotoxic compounds.

2173 Bioassays to Explore Mitochondrial Functions and Anticipate Drug Toxicity.

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The mitochondria have been identified as one of the major players in previously unrecognized drug-induced hepatic injuries. Hepatotoxicity is the main reason for drug withdrawal in clinical phases and post-market. Identifying early mitochondrial toxicities is thus essential to increase the chances of success in drug development. We have developed a panel of integrated functional bioassays aiming at measuring the different functions of the mitochondrial possibly associated with mitochondrial toxicity in a high throughput and robust manner. Three major axes were explored using cell-based bioassays: the bioenergetics, the redox status and the mitochondrial DNA depletion. First, measurements of the dioxygen consumption, cellular ATP level, extracellular lactate release with regard to cell viability, were multiplexed using fluorescence and luminescence methods. The identification of mitochondrial liabilities associated to hepatotoxicity was largely improved compared to the literature. Secondly, considering that the redox status is associated to drug toxicity, we measured the modulation of the mitochondrial ROS production and antioxidant defenses in response to drugs. Finally, due to its proximity to the electron transport chain and its lack of histories, mitochondrial DNA constitutes a potential target for drug-induced damages. The mitochondrial DNA content was quantified using high throughput analysis. We performed all the bioassays on HepG2 cells. We tested a panel of drugs that have been marketed, known to be responsible or not for mitochondrial liabilities associated or not to liver and/or cardiac injuries. The tested compounds were classified according to mechanisms of toxicity we detected and compared to the literature. Mitochondrial liabilities can be identified with a predicitivity of 84%. Coupling early indicators of mitochondrial dysfunction to late predictors of cytotoxicity provides more relevant and complete information on drug-induced mitochondrial injuries capable of reducing the large space of false negative left by existing methodologies.

2174 Glucose-Induced Gene Expression Changes Influence Barrier Function in Human Retinal Pigment Epithelial (RPE) Cells.

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The RPE forms a blood-retinal barrier in the back of the eye, and alterations in this barrier can adversely affect vision. In the present work, ARPE-19 cells were cultured in physiologically-relevant glucose levels (5 mM) and high glucose conditions (25 mM) for a period up to 3 weeks. Permeability, assessed by apical-basolateral flux of FITC-labeled dextran, was 10 ± 1 ng/ml/min in normal glucose and 6 ± 0.3 ng/ml/min in high glucose, suggesting an increase in barrier function. High glucose also increased mRNA levels of pigment-epithelial-derived factor (PEDF) about 5-fold, while decreasing VEGF-A mRNA levels about 2-fold; these changes correlated with secreted levels of these growth factors, which increased 2.3-fold and decreased 2.8-fold, respectively, in culture medium. High glucose levels increased expression of sphinogosine-1-phosphate receptor 1 (S1P1) in ARPE-19 cells more than 3-fold within 1 week of treatment, a change that was maintained during 3 weeks of exposure, whereas expression of S1P3 decreased in a time-dependent manner to nearly 7-fold lower levels after 3 weeks. mRNA levels of tight junction proteins zonula occludens-1 and claudin-1 were unchanged after 1 week of high glucose conditions, but decreased at 3 weeks by 2- and 8-fold respectively. The results of the present study indicate that, in ARPE-19 cells, high glucose conditions affect transcellular permeability and alter expression of proteins involved in tight junction regulation. Moreover, S1P1 and S1P3 genes, which are known to play critical roles in regulating the integrity of endothelial cell barrier function, may also contribute to the regulation of retinal barrier function in RPE cells.

2175 Towards an Integrated Risk Assessment of Hepatotoxic Drugs via Covalent Binding and Cellular Stress in Primary Hepatocytes.

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Introduction
Idiosyncratic DILI is of concern to drug development as it occurs rarely but with severe outcome. Early biomarkers for DILI besides covalent binding (CBV) are warranted to classify problematic drugs in vitro. Here we present on complementarity endpoints in primary hepatocytes that may early identify cellular damage.

Methods
Primary hepatocytes were treated with DILI drugs under attenuated oxidative stress via enzymatic in situ generation of H2O2 up to 24 h. CBV was determined by 14C binding, prostaglandins via quantification by LC-MS/MS in addition to ATP and LDH release. Expression profiles of Nrf2 regulated genes were created by qPCR analysis.

Preliminary Results
In hepatocytes generating H2O2, CBV was significantly exaggerated (13-fold for troglitazone; 20-fold for diclofenac) as compared to controls indicating further peroxisome activation of initially formed reactive metabolites. Significant increase of prostaglandin isomers was observed at lower drug concentrations and at earlier time points than changes in ATP and LDH. In rat hepatocytes prostaglandins E2 /
The objectives of this study were to: 1) determine the predictivity of the HC-MNT assay with the in vitro and in vivo MN assays for JAK2 inhibition imparts therapeutic benefit in inflammatory diseases, in particular rheumatoid arthritis. A novel series of JAK1-selective molecules was discovered using high-content micronucleus test (HC-MNT) in CHO cells. The current study we selected >50 compounds with known human hepatic or cardiac toxic parameters we could differentiate compounds with different organ toxicities. In the current study we selected >50 compounds with known human hepatic or cardiac toxicity. These were tested in both H9C2 cells (cardiac panel) and primary rat hepatocytes (hepatic panel) using high content mechanism screening. We found that the majority of compounds, regardless of their designated organ toxicities, displayed similar effects on the two panels tested. Only a small number of compounds demonstrated differential activity in the two panels (eg. nimesulide was only toxic to the hepatic panel). Our results indicate that sophisticated high content mechanism screening performed in an organ relevant cell line to assess a variety of mechanistic toxicity parameters we have recently shown that general cytotoxicity screening such as ATP measurement in organ specific cell lines can not accurately predict specific organ toxicity. However, we wanted to understand why by choosing more mechanistic parameters we could differentiate compounds with different organ toxicities. In both strains, hydroxyl DCF as well as DAG were detected as the main intestinal metabolites are excreted by the PCLS of TR- rats, but whether this is due to lower metabolism or increased permeability is not clear. However, it is not clear what are the direct consequences of Mrp2 deficiency in the intestine itself. Previously we reported that DCF was toxic in the rat intestine in vitro without the presence of liver metabolism. Therefore, using precision cut intestinal slices (PCIS), we compared wild type (WT) and Mrp2 deficient (TR-) rat intestine toxicity in vitro, by studying direct toxicity, DCF disposition and intracellular glutathione concentration. PCIS from WT and TR- rats were incubated with a concentration range of DCF, DCF induced similar dose-dependent toxicity and 200 μM DCF caused a significant decrease of ATP in both strains of rats indicating that the intestine from TR-rat is not intrinsically less sensitive to DCF toxicity. As glutathione is a substrate of Mrp2, Mrp2 deficiency may influence its accumulation and thereby the DCF induced toxicity. Intestinal GSH level in the TR-rats was significantly lower than in WT rats but did not make the TR- rat intestine more vulnerable. In both strains, hydroxyl DCF as well as DAG were detected as the main intestinal metabolites after 5 hours incubation, less amount was excreted into the medium by PCLS of the TR-rats. The study of DCF disposition is ongoing with using chamber. The result of drug safety remains a serious problem for the pharmaceutical industry. Extensive efforts are made to develop early predictive in vitro screens to assist in selecting compounds with a more desirable safety profile. Since cardiac and hepatic toxicity remain the top causes of compound attrition, much focus has been on building predictive assays for such toxicities. One platform is high content screening performed in an organ relevant cell line to assess a variety of mechanistic toxicity parameters. We have recently shown that general cytotoxicity screening such as ATP measurement in organ specific cell lines can not accurately predict specific organ toxicity. However, it is not clear what are the direct consequences of Mrp2 deficiency in the intestine itself. Previously we reported that DCF was toxic in the rat intestine in vitro without the presence of liver metabolism. Therefore, using precision cut intestinal slices (PCIS), we compared wild type (WT) and Mrp2 deficient (TR-) rat intestine toxicity in vitro, by studying direct toxicity, DCF disposition and intracellular glutathione concentration. PCIS from WT and TR- rats were incubated with a concentration range of DCF, DCF induced similar dose-dependent toxicity and 200 μM DCF caused a significant decrease of ATP in both strains of rats indicating that the intestine from TR-rat is not intrinsically less sensitive to DCF toxicity. As glutathione is a substrate of Mrp2, Mrp2 deficiency may influence its accumulation and thereby the DCF induced toxicity. Intestinal GSH level in the TR-rats was significantly lower than in WT rats but did not make the TR- rat intestine more vulnerable.

In conclusion, the TR- rat intestine is not intrinsically less sensitive to DCF toxicity and the lower GSH level does not make it more vulnerable. Less intestinal metabolites are excreted by the PCLS of the TR-rats, but whether this is due to lower production or lower excretion ability needs to be further validated.

The presented methods are useful for an early identification of DILI liabilities. Besides CVB, cellular stress in hepatocytes may contribute to development of DILI; here prostaglandins are sensitive biomarkers. mRNA levels give insight into the mechanism of toxicity. The in situ generation of H2O2 provides a supportive tool by attenuating response that is specific to drug treatment.

15(B) D2 increased by 3.3 / 2.3-fold for FeNTA and 3.0 / 2.0-fold for troglitazone at 6 h. In human hepatocytes, this increase was 3.6 and 3.8-fold respectively for troglitazone with no effect on LDH or ATP. mRNA levels indicate the activation of several keap/Nrf2 regulated genes such as NAD(P)H dehydrogenase (7,7-fold) and GSH-S-transferase ε2 (6.1-fold) after 24 h for troglitazone. Heme oxygenase 1 (6.4-fold) as well as GSH-S-transferase π1 (12.3-fold) showed a transient induction after treatment with FeNTA.

Conclusions

The presented methods are useful for an early identification of DILI liabilities. Besides CVB, cellular stress in hepatocytes may contribute to development of DILI; here prostaglandins are sensitive biomarkers. mRNA levels give insight into the mechanism of toxicity. The in situ generation of H2O2 provides a supportive tool by attenuating response that is specific to drug treatment.

The Rho GTPase Rac family are intracellular signaling proteins that control gene expression and various cellular functions including invasion and metastasis, cell cycle progression and apoptosis. Furthermore, they have been reported to be implicated in cancer initiation and progression. Rac1 is a member of the Rho family GTPases associated with lamellipodia or invadopodia causing invading cells to migrate, and is activated via association with Guanine Exchange Factors (GEFs), among which Vav2. Increase Rac1 activation has been associated with increased breast and brain cancer cell proliferation and invasion. Therefore, one main goal is to focus on the design of novel Rac inhibitors for the development of anticancer drugs. Previously, our laboratory synthesized EHop-016, which was demonstrated to be the first known inhibitor of Vav2-Rac1 interaction in MDA-MB-435 metastatic cancer cells at low micromolar concentrations. In order to reduce the toxicity and increase the potency, we utilized molecular docking to design novel EHop-016 derivatives. The carbazole group of EHop-016 appeared to be required for inhibitory activity, and thus was maintained as a core fragment in further design. It appeared from the docking experiments that replacement of the central pyrimidine ring with other building blocks that orient the potential inhibitors into a U-shaped conformation, provided the best docking results. Novel molecules, that according to docking results bind much better to Rac1 than EHop-016 will be presented, and the specific interactions leading to increased binding will be discussed. As the compounds were designed to be easily accessible via laboratory synthesis, these proposed improved inhibitors of Rac activity could lead to novel antitumorigenic cancer therapies.

C. Racing, T. Ferguson, S. Bakoi, D. Preston, D. Anestis and G. Rankin. Pharmacology, Physiology and Toxicology, Marshall University, Huntington, WV.

Chlorinated anilines are common intermediates in the production of agricultural chemicals, dyes, industrial compounds, and pharmaceuticals. Some chloroanilines can induce nephrotoxicity in vivo and in vitro. Previous studies have shown 3,5-dichloroaniline (3,5-DCA, 1.0 mM) induced nephrotoxicity in isolated renal cortical cells (IRC) following 90 min exposure. Studies from our lab have also shown IRCC pretreated with non-selective cytochrome P450 (CYP) inhibitors [piperonyl butoxide (1.0 mM) and metyrapone (1.0 mM)] partially attenuated 3,5-DCA toxicity, suggesting that CYPs may play a role in 3,5-DCA bioactivation. The purpose of the present study was to further explore the role of CYP mediated 3,5-DCA bioactivation using an in vitro rat model. IRCC were obtained from male Fischer 344 rats. IRCC (4 x 106 cells/ml; 3mL) were incubated with shaking for 90 min with either dimethyl sulfoxide (DMSO) or 3,5-DCA (1.0mM). IRCC were pretreated with various CYP inhibitors [isoniazid (1.0 mM), ketoconazole (0.1 mM), omeprazole (0.01 mM), diethylthiocarbamate (DEDTCA; 0.1 mM), oleandomycin triacetate (0.5 mM), or sulfaphenazole (0.1 mM)] and cytotoxicity was determined by measuring lactate dehydrogenase (LDH) release. Pretreatment with DEDTCA, omeprazole, and sulfaphenazole partially attenuated 3,5-DCA induced nephrotoxicity, while ketomiconzole (0.01 mM), oleandomycin triacetate (0.5 mM) also did not alter 3,5-DCA induced nephrotoxicity. Studies in rats have shown that DEDTCA, omeprazole, and sulfaphenazole are effective inhibitors of the CYP2C family isozymes. These results suggest that 3,5-DCA is bioactivated via multiple pathways, one of which involves the CYP2C family. (Supported in part by NIH Grant 8F20GM103434 to the West Virginia IDEA Network for Biomedical Research Excellence)


Serum creatinine level is a commonly used surrogate measure for glomerular damage and kidney function evaluation. It is well known that creatinine clearance in the kidney is through free glomerular filtration and transporter mediated excretion in proximal tubule. Inhibiting these transporters can result in transient increase in plasma creatinine level, which can lead to false interpretation of glomerular damage.

Recently, we have identified that, in addition to OCT2 and MATE1, OAT2 and OAT3 can play major role on creatinine secretion. This finding led to development of a novel cellular model co-expressing all four transporters, for modeling active creatinine secretion in the kidney.

Using the model, we tested drugs with reported incidence of increasing serum creatinine. Most of these drugs inhibited B>A tranacellular transport of [C14]Creatine to different extent, suggesting they are able to block active tubular secretion of creatinine in vivo. It is noteworthy that Citomizedine, which has been previously demonstrated by us as a pan-inhibitor of all four transporters, was able to completely abolish creatinine transport; whereas trimethoprimer, which is not an OAT2 inhibitor, only resulted in partial inhibition. These results are in good accordance with clinical studies that higher dosage of Citomizedine can completely block active creatinine secretion in proximal tubule whereas there is no statistically difference in serum creatinine level in patients under moderate- and high-dose of trimethoprimer. Furthermore, salicylic acid, which was tested at clinically relevant concentrations, also exhibited a dose-dependent and substantial inhibitory effect on active creatinine secretion. Overall these results suggest that in addition to other drugs, the possible contributing factors to serum creatinine level increase caused by salicylic acid, possibly through inhibiting OAT2 and/or MATEs transporters.


The Rho GTPase Rac family are intracellular signaling proteins that control gene expression and various cellular functions including invasion and metastasis, cell cycle progression and apoptosis. Furthermore, they have been reported to be implicated in cancer initiation and progression. Rac1 is a member of the Rho family GTPases associated with lamellipodia or invadopodia causing invading cells to migrate, and is activated via association with Guanine Exchange Factors (GEFs), among which Vav2. Increase Rac1 activation has been associated with increased breast and brain cancer cell proliferation and invasion. Therefore, one main goal is to focus on the design of novel Rac inhibitors for the development of anticancer drugs. Previously, our laboratory synthesized EHop-016, which was demonstrated to be the first known inhibitor of Vav2-Rac1 interaction in MDA-MB-435 metastatic cancer cells at low micromolar concentrations. In order to reduce the toxicity and increase the potency, we utilized molecular docking to design novel EHop-016 derivatives. The carbazole group of EHop-016 appeared to be required for inhibitory activity, and thus was maintained as a core fragment in further design. It appeared from the docking experiments that replacement of the central pyrimidine ring with other building blocks that orient the potential inhibitors into a U-shaped conformation, provided the best docking results. Novel molecules, that according to docking results bind much better to Rac1 than EHop-016 will be presented, and the specific interactions leading to increased binding will be discussed. As the compounds were designed to be easily accessible via laboratory synthesis, these proposed improved inhibitors of Rac activity could lead to novel antitumorigenic cancer therapies.

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bolic transcription factors nuclear factor κB (NF-κB) and signal transducer and activator of transcription factor-3 (STAT-3), respectively, were all increased in STZ-rats and suramin blocked these changes. Therefore, delayed administration of suramin attenuated urinary markers of DN, inflammation by blocking NF-κB activation/ICAM-1-mediated leukocyte infiltration and fibrosis by blocking STAT-3 -TGF-β1/Smad-3 signaling, supporting the potential use of suramin in development of diabetes-induced kidney injury.

2185 Resveratrol Alterns Subcellular Changes in Oxidative Stress Mediated by Cisplatin in Rat Renal Tissue.
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The cancer chemotherapeutic agent cisplatin is associated with a 33% incidence of diminished renal function. Interventions to reduce renal toxicity are important in improving patient outcome. Resveratrol (RES) is a phytochemical found in grapes, cranberries and nuts. RES has been recognized as a natural agent possessing anti-cancer and antioxidant properties. This study investigated RES attenuation of cisplatin renal in vitro toxicity and focused on differences between cytosolic and mitochondrial oxidative stress mediated by cisplatin. Male Fischer 344 rats (200-250 g) were anesthetized, with isoflurane and the kidneys were isolated. Renal cortical slices were prepared and pre-incubated with 30 ul ethanol (VEH) or 30 ug/ml resveratrol (RES, final concentration) for 30 min at 37°C. Tissue was subsequently incubated for a maximum of 120 min with 0.7%, or 150 ug/mL cisplatin. Loss of membrane integrity was evaluated as leakage of lactate dehydrogenase (LDH). Oxidative stress and nitrosative stress was assessed in kidney homogenate as well as in subcellular fractions of mitochondria and cytosol. Oxidative stress was measured by protein carbonylation using an Oxyblot. Nitrosative alterations were assessed using 3-nitrotyrosine formation by western blot. LDH leakage required a 120 min exposure to cisplatin. An increase in protein carbonyls as detected by Oxyblot, was increased by cisplatin and totally prevented by RES. RES also decreased protein carbonyls in tissue not exposed to cisplatin. Our findings showed that a 30 min RES pre-incubation diminished cisplatin renal toxicity and that protein carbonyl and 3-nitrotyrosine and early changes in oxidative stress prior to the onset of loss of membrane integrity. (Supported by NIH Grants INBRE 5P20RR016477-09S4; 5P20RR016477 and 8P20GM103434 to the West Virginia IDEA Network for Biomedical Research Evaluation).

2186 Diglycolic Acid Induces Cytotoxicity in Human Proximal Tubule Cells via Preferential Inhibition of Sucinate Dehydrogenase and Oxidative Phosphorylation.
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Diethylene glycol (DEG) is an organic solvent used in common consumer products, thus allowing for increased risk for exposure. DEG metabolism produces two primary metabolites, 2-hydroxyxethyacetic acid (2-HEAA) and diglycolic acid (DGA). DGA, not DEG or 2-HEAA, produces proximal tubule cell necrosis leading to acute renal failure, the hallmark of DEG poisoning. Studies were designed to assess whether the mechanism for DGA-induced cytotoxicity involves disruption of cellular metabolic processes resulting in mitochondrial dysfunction. DGA induces severe ATP depletion in human proximal tubule (HPT) cells that occurs prior to significant cell death. HPT cells pretreated with increasing DGA concentrations showed significant decreases in oxygen consumption suggesting that DGA acts as an oxidative phosphorylation inhibitor, rather than a mitochondrial electron transport chain uncoupler. Co-incubation of DGA with the antioxidant, t-tocopherol significantly reduced DGA-induced reactive oxygen species (ROS) formation, but did not reduce ethidium homodimer uptake or lactate dehydrogenase release, two indicators of apoptosis. Oxidative stress, as measured by protein carbonyl and 3-nitrotyrosine formation, and early changes in oxidative stress prior to the onset of loss of membrane integrity were significantly reduced DGA-induced reactive oxygen species (ROS) formation, but did not reduce ethidium homodimer uptake or lactate dehydrogenase release, two indicators of apoptosis. Oxidative stress, as measured by protein carbonyl and 3-nitrotyrosine formation, and early changes in oxidative stress prior to the onset of loss of membrane integrity were significantly decreased by DGA treatment. MTT staining in LNCaP cells, but neither TSA nor 5-Aza altered MTT staining. Treatment of prostate cancer cells with CBZ prior to docetaxel exposure decreased MTT staining in LNCaP cells, but neither TSA nor 5-Aza altered MTT staining compared to cells exposed to docetaxel alone. Treatment of normal rat kidney (NRK) and human embryonic kidney 293 (HEK293) cells with the same concentration of epigenetic inhibitors used in prostate cancer cells significantly decreased MTT staining after 48 hr. In contrast, a frontline chemotherapeutic, did induce concentration-dependent decreases in MTT staining after just 24 hr. Treatment of prostate cancer cells with CBZ prior to docetaxel exposure decreased MTT staining in LNCaP cells, but neither TSA nor 5-Aza altered MTT staining compared to cells exposed to docetaxel alone. Treatment of normal rat kidney (NRK) and human embryonic kidney 293 (HEK293) cells with the same concentration of epigenetic inhibitors used in prostate cancer cells significantly decreased MTT staining after 48 hr. The epigenetic inhibitors were generally more toxic than the xenobiotic bromate (0-400 ppm), and TSA and 5-Aza pretreatment increased the cytotoxicity of BrO3- in NRK cells. Increased cytotoxicity in NRK cells correlated to alterations in epigenetic markers such as histone phosphorylation and to alterations in the methylation of the cyclin-dependent-kinase inhibitor protein p21. Collectively, these data suggest that epigenetic inhibitors can induce nephrotoxicity at doses that are therapeutically relevant in prostate cancer cells In Vitro.

2187 Chronic Ethanol Ingestion in Mice Induces Renal Inflammation and Injury through the Platelet-Activating Factor Receptor.
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Ethanol exposure increases circulating oxidized phospholipids in rats and in alcoholic steatopathisis patients, although the biological role of circulating oxidized phospholipids remains unclear. Platelet-activating factor (PAF), and some oxidized phospholipids, activate the Platelet-activating factor receptor (PAFR). We hypothesized that PAFR contributes to renal inflammation and injury after ethanol ingestion. In this study we find that mice fed a Lieber-DeCarli ethanol diet had increased levels of circulating PAF, and the pro-apoptotic oxidatively truncated phospholipid azelaoyl-PC. Most strikingly, PAFR-null mice had less infiltrating neutrophils, oxidized phospholipids and TUNEL positive cells in kidney as compared to wild-type (wt) mice fed with ethanol diet. Blood urea nitrogen and serum creatinine, renal functional markers, increased after ethanol exposure in wt mice but not in PAFR-null mice. The renal fibrosis marker, α-smooth muscle actin was increased in ethanol-fed wt mice, but not in PAFR-null mice. Kidney injury molecule-1 (KIM-1), a marker for proximal tubular injury, is increased during acute renal injury, and we find higher expression of KIM-1 in the proximal tubules of ethanol fed wt mice as compared to PAFR-null mice. These results suggest that PAF and bioactive PAF-like oxidized phospholipids promote renal inflammation, acute injury, and renal fibrosis. These events completely depend on a functional PAFR. This demonstrates a novel mechanism of PAFR in alcoholic renal injury.

2188 Nephrotoxicity of Epigenetic Inhibitors Used for the Treatment of Cancer.
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Pharmacological and Biomedical Sciences, University of Georgia, Athens, GA.

Several studies exist investigating the anti-neoplastic activity of epigenetic inhibitors. In contrast, fewer studies have investigated the toxicity of epigenetic inhibitors in non-target organs, such as the kidney. Even fewer have investigated both the anti-neoplastic activity and toxicity of epigenetic inhibitors under similar conditions. This study determined the anti-neoplastic activity and nephrotoxicity of epigenetic inhibitors In Vitro. The therapeutic efficacy of epigenetic inhibitors was first determined in human prostate cancer cells (PC-3 and LNCaP) using the DNA methyltransferase inhibitor 5-aza-cytidine (5-Aza) and the histone deacetylase inhibitor trichostatin A (TSA). Cells were also treated with carbamazepine (CBZ), an anti-convulsant with histone deacetylase inhibitor-like properties. 5-Aza, TSA, or CBZ alone (0-100 μM) did not induce decreases in MTT staining in PC-3 or LNCaP cells after 48 hr. In contrast, a frontline chemotherapeutic, did induce concentration-dependent decreases in MTT staining after just 24 hr. Treatment of prostate cancer cells with CBZ prior to docetaxel exposure decreased MTT staining in LNCaP cells, but neither TSA nor 5-Aza altered MTT staining compared to cells exposed to docetaxel alone. Treatment of normal rat kidney (NRK) and human embryonic kidney 293 (HEK293) cells with the same concentration of epigenetic inhibitors used in prostate cancer cells significantly decreased MTT staining after 48 hr. The epigenetic inhibitors were generally more toxic than the xenobiotic bromate (0-400 ppm), and TSA and 5-Aza pretreatment increased the cytotoxicity of BrO3- in NRK cells. Increased cytotoxicity in NRK cells correlated to alterations in epigenetic markers such as histone phosphorylation and to alterations in the methylation of the cyclin-dependent-kinase inhibitor protein p21. Collectively, these data suggest that epigenetic inhibitors can induce nephrotoxicity at doses that are therapeutically relevant in prostate cancer cells In Vitro.

2189 hTERT-Immortalized Renal Proximal Tubule Epithelial Cells: A Model for Testing Cadium’s Role in the Development of Renal Cancer.
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Global Environmental Health Sciences, Tulane University, New Orleans, LA. Sponsor: J. Wickliffe.

The incidence of renal cell carcinoma (RCC) has steadily increased in the United States over the past three decades. Therefore, it is critical that scientists are better able to understand contributing factors. Cadmium (Cd) is a recognized carcinogen and widespread environmental contaminant. The kidney is a known target organ of Cd toxicity; however the mechanistic role(s) that Cd plays in the development of RCC has not been described. Renal proximal tubule epithelial cells (RPTECs) are specifically affected by Cd because of their propensity to reabsorb Cd from filtrate.
leading to bioaccumulation. In order to elucidate the mechanisms by which Cd acts, we aim characterization of newly developed cell line derived from renal proximal tubule epithelial cells of a healthy human male donor (RPTEC/TERT1). Our goal is to establish this new in vitro system for future toxicological and cancer research by characterizing the RPTEC/TERT1 cell line and utilizing them to conduct biologically relevant exposure studies. We will explore these goals through exposure experiments to binary mixtures of two common environmental contaminants, Cd and benzo(a)pyrene (B(a)P). Our preliminary results demonstrate that these cells are sensitive to both compounds separately over a wide range of exposure levels. Additionally, the characteristic Cd-interacting protein, metallothionein I/II, exhibits a Cd-specific protein response, which supports its ability as a function of exposure, based on in vivo evidence. The RPTEC/TERT1 cell line expresses metabolic biotransformation enzymes necessary for processing polycyclic aromatic hydrocarbons (PAHs) which is exemplified by their sensitivity to the representative PAH, B(a)P. Future experiments will further characterize DNA repair capacity of the RPTEC/TERT1 cell line and development of mutagenic lesions when exposed to binary mixtures of sub-cytotoxic doses of Cd and B(a)P. This work will provide mechanistic support and augment current scientific knowledge regarding the development of RCC and exposure to xenobiotic mutagens and carcinogens.

**2190 BUN and Serum CRE Alterations in Higa and BALB/c Mice after Subacut Administration of Fluoride via Drinking Water.**

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Fluoride (F) is known as an environmental pollutant. Because F is filtered by the kidney, mice with impaired renal function may be affected more significantly by F. IgA nephritis is the most common chronic glomerulonephritis. High IgA (HGA) mice have been used as a model of IgA nephritis. The effects of fluoride on BUN and serum creatinine (CRE) of HGA and BALB/c mice after subacute administration via drinking water were examined in this study to get basic information of the toxic effects of F on the mice with IgA nephritis. F was administered to HGA and BALB/c mice, aged 11-12 weeks at 0, 50, 100, and 150 ppm in their drinking water for 4 weeks. The blood was sampled from the tail artery once a week. The BUN and CRE in the serum were determined by the kits. For the BUN levels in the HGA mice, after one week from the beginning of the exposures, the mean BUN in the 50-ppm group was significantly higher than those in the 100- and 150-ppm groups. At 2 weeks, the mean value of BUN in the 100-ppm group was significantly higher than those in the 0- and 50-ppm groups. For the BALB/c mice, there were no significant differences among the groups. For the CRE in the serum of HGA mice, after 1 week, the mean CRE in the 50-ppm group was significantly higher than those in the 100- and 150-ppm groups. For the BALB/c mice, after 3 weeks, the mean CRE in the 50-ppm group was significantly higher than those in the 100- and 150-ppm groups. The alterations in the BUN and CRE in the serum observed in the HGA and BALB/c mice may indicate the toxic effects of F on the kidney. However, the alterations were not dose-dependent, and the effects of F on the kidney of HGA mice were not as significant as those on kidney of the BALB/c mice. The toxic effects of F on the kidney were not enhanced in the HGA mice at 11 to 12 weeks of age.

**2192 Miniaturized Multiplex Protein NANOARRAYS For Early Detection of Drug-induced Nephrotoxicity in Mice and Rats.**


Drug-induced toxicity is the major cause of kidney damage. The previous methods for measuring drug-induced injury, namely serum creatinine and blood urea nitrogen, have often been found inadequate. Recently the FDA and EMEA have qualified a panel of biomarkers that is highly useful in the early identification and characterization of kidney injury. Here we report the development of highly sensitive and small sample volume NanoArray multiplex assays for early detection of drug-induced nephrotoxicity in mice and rat in-vivo models based on dip-pen nanolithography (DPN™) using a NanoArrayer 3000TM. This instrument enables the production of highly reproducible micro- to nano-scale arrays of proteins which occupy an area approximately 100 times smaller than conventional microarrays. The reduced area results in reduced reagent and sample requirement and increased sensitivity due to reduced analyte depletion. For a multiplex sandwich ELISA, arrays of specific capture antibodies are printed on activated glass slides. Samples of as little as 2-4 μl can be incubated over the arrays in 48 and 96 well format. The slides are then incubated with specific biotinylated detection antibodies followed by fluorescently labeled Streptavidin. The fluorescence is measured with a high resolution fluorescence scanner. The multiplex assays were developed and validated for mouse (Albumin, Clusterin, Cystatin C, KIM-1, and EGF) and rat biomarkers (Albumin, Clusterin, Cystatin C, KIM-1, B2MG and TFF3) and showed high sensitivity and specificity. The assay was used to detect and quantify renal injury biomarkers in urine samples obtained from normal and drug compromised rats. The low volume sample requirements in conjunction with the extremely high sensitivity, selectivity, and reproducibility help enable longitudinal studies on individual rodents thus saving significant study costs and providing better toxicity data than other conventional methods used for preclinical and clinical safety and toxicity studies.

**2193 Magnetic Resonance Imaging (MRI) Assessment of Renal Glomerular Filtration Rate (GFR).**

M. Uteng1, A. Mah1, A. Piaia2, J. Cuncliffe3, E. Persohn4, E. Trinoto5, D. Ledieu1, P. Moulin1, N. Shangari1, S. Chibout1, A. Wolf1, L. Li1, E. Pogman1 and N. Beckmann2. 1Novartis Institutes for BioMedical Research, Novartis, Basel, Switzerland; 2Global Imaging Group, Novartis, Basel, Switzerland.

MRI of kidneys after administration of the contrast agent, gadodiamide-tetra-aza-clo-dodecanetetra-acetic acid (Gd-DOTA), has been reported as a promising method for assessment of GFR. However, there is little literature on the sensitivity of this method compared to standard assessments in non-clinical trials of drug development. To this end, we have conducted a study with the aim to determine the sensitivity of MRI as a method for detection of GFR in rodents. Sprague-Dawley rats were treated with Adefovir as positive control, and Telbivudine and Entecavir as negative controls for nephrotoxicity. The rats were treated daily by oral gavage for 4 weeks with 10X and 25X human equivalent exposure doses (HED) of compounds. After 3, 10 and 28 days treatment, GFR was assessed by MRI and creatinine clearance rate, while renal toxicity was assessed by urinary kidney biomarkers, clinical biochemistry parameters in blood and urine, histopathology, electron microscopy and gene expression profiling. Impairment of GFR was detected by MRI showing delayed time-to-peak signal of Gd-DOTA clearance in the renal cortex of animals treated with 25X HED Adefovir nanocrystal resuspension mixture. The linear correlation between GFR and Gd-DOTA clearance was excellent. MRI was more sensitive in detecting GFR impairment than urinary kidney biomarkers, clinical biochemical parameters in blood and urine, histopathology, electron microscopy and gene expression profiling. Furthermore, we have demonstrated the potential of MRI as a non-invasive, longitudinal method for assessment of GFR in preclinical and clinical trials of drug development.

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genes. On the other hand, the creatinine filtration rate did not reveal any significant changes. Cells treated with Trehalose and Entecavir showed neither renal impairment nor toxicity for the entire study duration.

In conclusion, these results demonstrated that MRI is a sensitive method for non-invasive detection of GFR changes and that this technique may be a good alternative to the standard measurements of creatinine filtration in non-clinical investigative studies.

**2194 Identification of 3-Indoxyl Sulfate As an Early Biomarker for Nephrotoxicant-Induced Acute Kidney Injury.**

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The identification of new biomarkers of acute kidney injury (AKI) is important for the detection of drug-induced kidney damage. Various serum or urinary biomarkers have been used to detect AKI, but these biomarkers have shown poor sensitivity and specificity. In this study, we compared the sensitivity of a new metabolomic biomarker, 3-indoxyl sulfate (3-IS), with traditional biomarkers for the diagnosis of AKI using the area under the receiver operating characteristic (ROC) curve. Sprague-Dawley male rats were allocated to several groups. Each group was administered either a single dose of cisplatin (20 mg/kg, i.p.), continuous injection of cyclosporin A (10 mg/kg, i.p.), mercury chloride (1.5 mg/kg, i.p., and 7.5 mg/kg, i.p.), or gentamicin (60 mg/kg, s.c.). Urine and plasma samples were collected 1, 3, and 7 days after last injection of nephrotoxicants. Urine and blood biochemical parameters involved in kidney toxicity were measured. We also measured 3-IS levels in the serum, urine, and kidney using HPLC. In the nephrotoxicants-treated rats, blood urea nitrogen (BUN) and serum creatinine (sCr) levels were slightly increased. The 3-IS levels were significantly reduced in the urine of rats treated with cisplatin and other nephrotoxicants. In contrast, 3-IS levels were significantly elevated in the urine and kidneys of nephrotoxicants-treated rats. The 3-IS is produced by bacterial metabolism of tryptophan in the intestine, followed by oxidation and transport in the liver. The 3-IS is mainly excreted via the urinary tract via the organic anion transporter (OAT) in the proximal tubule. Therefore, reduced urinary 3-IS levels can reflect proximal tubule injury. These results suggest that urinary 3-IS may be used as an alternative to traditional biomarkers to predict AKI.

**2195 A Quantitative High-Throughput Screening Platform for Predictive Kidney Toxicity.**

M. Adler1, E. Gottwald1, B. Goodwin2, M. Xia2, and V. S. Vaidya2. 1Renal Division, Department of Medicine, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA; 2National Center for Advancing Translational Sciences, National Institutes of Health, Bethesda, MD.

Drug and environmental chemical-induced kidney toxicity plays an important role in the high incidence and prevalence of kidney injury in both hospitalized and non-hospitalized patients, which in many circumstances can be prevented or at least minimized by predictive toxicity screening. The goal of this study was to develop a cell-based quantitative high throughput screening (qHTS) platform with two aims: 1) to identify a more biologically relevant in vitro system for prediction of human kidney toxicity than currently used immortalized cells and 2) to identify a sensitive, specific, robust and translatable biomarker of kidney toxicity since in vivo biomarkers such as kidney injury molecule-1 do not respond in vitro. We used primary human proximal tubular epithelial cells (HPTEC) and observed that these cells in a monolayer possess human tubular epithelial characteristics like 1) formation of domes; 2) expression of zonula occludens-1, cytokeratin 18 by immunostainings; 3) a wide range of eflux and influx transporters including aquaporin 1, megalin, or domes; 2) expression of zonula occludens-1, cytokeratin 18 by immunostainings; 3) a wide range of eflux and influx transporters including aquaporin 1, megalin, organic cation transporter 2, multidrug resistance protein 2, P-glycoprotein by semi-quantitative PCR and 4) activity of brush-border enzymes like alkaline phosphatase, and γ-glutamyl-transferase. We found that hemoxgenease-1 (HO-1) mRNA and protein levels significantly increased in a concentration-dependent manner (tested over 6-point concentration curve) following exposure to structurally and mechanistically diverse kidney toxicants such as cisplatin, gentamicin, cyclosporin A and cadmium chloride and correlated well with cytotoxicity. HO-1 expression remained unchanged following treatment of HPTEC with non-kidney toxic compounds (e.g. carboplatin), demonstrating its specificity. Our results demonstrate the relevance and potential use of HPTEC in a qHTS platform using HO-1 as a biomarker for predictive safety assessment of drugs and environmental chemicals.

**2196 Systems Biology Approach Identifies Transcriptional Regulator of Kidney Injury Molecule-1.**

A. K. Avaj1, T. Kim1,3, P. J. Park2, V. Ramirez3, and V. S. Vaidya3. 1Renal Division, Brigham and Women’s Hospital, Boston, MA; 2Centre for Biomedical Informatics, Harvard Medical School, Boston, MA; 3Department of Medicine, Harvard Medical School, Boston, MA.

Kidney injury molecule-1 (KIM-1) is the highest upregulated gene following kidney ischemic or toxic insult and functions as a phosphatidylserine receptor to internalize apoptotic cells. Owing to lack of information about regulation of KIM-1, we used genome-wide expression data following kidney ischemia reperfusion injury (IRI) in rats and utilizing ChIP enrichment analysis and kinase enrichment analysis we identified STAT3 and checkpoint kinase 1 (Chk1) as a potential transcription factor and kinase regulating KIM-1. Then we performed an extensive biochemical validation of the bioinformatics predictions and report that reactive oxygen species generation following IRI upregulates Chk1 that binds to STAT3 phosphorylating it at Ser727, which further binds to KIM-1 promoter for its transcription.

We observed temporal association among pSTAT3, pChk1 and KIM-1 using immunoblotting and immunostaining in rat kidneys following IRI and in human kidneys from patients with kidney injury. To prove transcriptional regulation of KIM-1 by STAT3 we used i) primary human proximal tubular epithelial cells (HPTEC) and showed a significant increase (1.5 fold) in KIM-1 mRNA and protein following STAT3 activation. Conversely we used human renal carcinoma cell line (769 P) expressing high pSTAT3/KIM-1 and found 2-fold decrease in KIM-1 following STAT3 siRNA transfection. Furthermore, we confirmed that STAT3 binds on KIM-1 promoter in i) rat kidneys (10 fold by ChIP assay) following IRI; ii) HPTEC transfected with KIM-1-luciferase plasmid (3-fold by STAT3 activation) and iii) 769 P cells transfected with STAT3 siRNA (2-fold decrease). The binding of Chk1 to STAT3 was observed using immunoprecipitation in HPTEC by hydroxyurea (Chk1 activator). These results reveal Chk1-STAT3 as one of the key pathways regulating KIM-1 transcription.

**2197 Kidney miRNAs Show Age and Sex Differences in Expression during the Rat Life Cycle.**


Increasing evidence for epigenetic mechanisms of gene regulation has fueled interest in the role of miRNAs in toxicogenomics for biomarker discovery. While relatively immune in comparison to other genomic resources, the growing knowledge base of individual miRNAs and their putative gene targets allows for large scale inquiry into more comprehensive, genome-wide analysis of miRNA expression. Kidney tissues in the F-344 rat model system were examined over the life cycle for the purpose of evaluating miRNAs with putative roles in drug metabolism and kidney disease. miRNA expression was characterized at 2, 5, 6, 8, 15, 21, 78, and 104 weeks of age in both sexes using Agilent 8x15k rat miRNA microarrays containing multiple probes for 677 unique miRNAs. Five animals per sex and age were used for a total of 80 samples. Agilent’s Feature Extraction software was used for initial analysis and processing of the raw data and 224 miRNAs were found to be expressed in the kidney in at least one age and sex. Combined filtering criteria of 1.5 fold change and p < 0.05 (2-way ANOVA) revealed 105 miRNAs (47%) exhibiting differential expression by age or sex. Principal component analysis (PCA) showed PC1 accounted for 21% of the variability among the 105 differentially expressed miRNAs in a pattern consistent with age-specific effects. 12 miRNAs showed increased expression at 78 and 104 weeks, consistent with an aging-related effect (e.g. miR-142-3p, miR-223). Although no large scale, sex-related patterns were evident from the PCA, some miRNAs showed sex-specific patterns of expression (e.g. miR-204, miR-499, miR-183). miR-499 has been implicated in regulation of mitochondrial dynamics through direct targeting of calcineurin. Collectively, these results comprise one of the first large-scale characterizations of global miRNAs in the kidney over the entire rat life cycle and sex-related differences that may impact susceptibility to adverse effects in the kidney.

**2198 Mice Deficient in microRNA-155 Have Greater Susceptibility to Cisplatin-Induced Kidney Toxicity.**

K. L. Pellegrini, V. Bijol and V. S. Vaidya. Renal Division, Department of Medicine, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA.

Although originally identified as an oncogenic factor, microRNA-155 (miR-155) has also been found to be upregulated in macrophages and dendritic cells in response to a range of inflammatory stimuli and required for the activation of Th17 cells. We have previously shown miR-155 to be significantly upregulated following...
2199 Urinary microRNAs As Translational Biomarkers to Detect Acute Kidney Injury.
K. Ramachandran, J. Saikumar, S. S. Waikat and V. S. Vaidya. Renal Division, Department of Medicine, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA.

MicroRNAs (miRNAs) are a family of short, single stranded, non-coding RNA molecules that direct the expression of nearly 60% of all protein coding genes. Extracellular miRNAs, identified in 14 different body fluids, have been proposed as biomarkers of disease and organ damage due to their stability, sensitivity, specificity and ease of detection. Previously, we identified 3 miRNAs (miR-21, -155 and -18a) that were upregulated following ischemic injury and gentamicin induced nephrotoxicity in rats. Human urinary levels of miR-21 and -155 were able to distinguish patients with and without acute kidney injury. The aim of our current study was to profile the human miRNome in the urine of patients with or without AKI to identify a panel of urinary miRNAs that can serve as sensitive and specific indicators for kidney injury. To estimate the fraction of miRNAs present in the urine, we used the Human miRNome microscript miRNA PCR Array from Qiagen (miRBase version 18 containing ~1900 miRNAs) on urines pooled from 6 patients with AKI and 6 healthy controls. Samples were collected from patients admitted in the Intensive Care Unit (ICU) with a rise in serum creatinine of 100% over baseline. Using a cycle threshold (Ct) range of 19-30, miRNAs that were expressed in both, or in either one of the diseased or healthy pools were selected. All miRNAs that had Ct values >30 in both the sample sets were considered as ‘Not Expressed and eliminated. Thus, we designed a customized array of the 378 detected miRNAs and analyzed the 12 urine samples (6 AKI and 6 healthy) individually. We found 52 miRNAs that were upregulated >-5 fold in the AKI patients as compared to the controls and 33 candidate miRNAs (out of 52) were selected using a standard deviation cut-off of 1.5. Further evaluation of these candidate miRNAs for sensitivity, specificity, stability, reproducibility and robustness in an expanded cohort of patients with or without kidney damage will help in establishing the value of urinary miRNAs as non-invasive biomarkers for kidney injury.

2200 Genetic Reduction in Fibrinogen Protects from Progression of Acute Kidney Injury to Chronic Kidney Disease.
F. Craciun, A. K. Ajay and V. S. Vaidya. Department of Medicine, Renal Division, Brigham and Women’s Hospital, Harvard Medical School, Boston, MA.

The incidence of acute kidney injury (AKI) is increasing and recent studies emphasize the significantly elevated microRNA (miRNA) and mRNA levels in the kidney injury. We have shown that miRNA and protein expression in the kidney as well as urinary fibrinogen (Fg) is significantly increased following AKI in mice, rats and humans. Furthermore, we have demonstrated that Fg heterozygosity in mice reduces plasma Fg to 75% of the normal circulating levels, protecting from AKI and promoting faster resolution of kidney damage. Therefore, we hypothesized that Fg heterozygosity would protect from AKI to CKD progression. To test this we used a Folic acid (250 mg/kg, single sq injection) induced AKI to CKD progression model in wildtype (Fg+/+), heterozygous (Fg+/-) and Fg deficient (Fg-/-) mice on Balb/c background and the mice were sacrificed at days 1 and 14 following administration (n=4/time point/group). At day 1 there was significantly reduced kidney dysfunctions as assessed by blood urea nitrogen (112, 73 and 65 mg/dL for Fg+/+, Fg+/- and Fg-/- respectively) and serum creatinine (0.7, 0.6, 0.3 mg/dL for Fg+/+, Fg+/- and Fg-/- respectively), indicating protection in Fg+/- and Fg-/-.

Kidney mRNA levels of kidney injury molecule-1 (marker of proximal tubular injury) were significantly higher for Fg+/- (80 fold) than Fg+/- (46 fold) and Fg-/- (55 fold) as compared to unjured mice. By day 14 these markers of acute injury reverted to normal but there was increased kidney mRNA expression of fibrosis markers fibronectin (9, 4 and 8 fold in Fg+/+, Fg+/- and Fg-/- respectively) and collagens (8, 3 and 9 fold in Fg+/+, Fg+/- and Fg-/- respectively) when compared to unjured mice, with unjured Fg-/- showing protection. This was confirmed histologically by Masson’s trichrome staining. We conclude that a lowered but not completely abolished level of Fg protects from AKI to CKD progression, providing a therapeutic target that could benefit AKI survivors.

2201 Urinary Levels of N-Acetyl-β-D-Glucosaminidase (NAG), Glutathione-S-Transferase (GST), Blood Lead and Plasma Creatinine As Early Indicators of Lead Nephropathy in Occupationally Exposed Subjects.
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Lead (Pb) toxicity remains a public health problem. Although early diagnosis is paramount for meaningful intervention particularly in occupationally exposed subjects, the diagnosis of Pb poisoning at early stage remains a problem even in developed countries. This work addressed this issue by expressing urinary levels of NAG and GST as exponents of concentrations of conventional renal function markers (creatinine, urea and uric acid) and blood Pb. The result of exponential expression of these results showed a definite pattern particularly in occupationally exposed subjects relative to confirmed chronic renal failure subjects and control. When these figures were compared logarithmically, definite hyperboles for the control, the occupationally exposed and the CRF were observed; the importance of this in early diagnosis of lead nephropathy is to be discussed. Based on these, we propose hypothesis that can be used in the early identification of renal tubular damage especially in subjects occupationally exposed to lead using this logarithmic model.

2202 Early Postnatal Gentamicin Treatment Reduces Glomerular Number in Extra Uterine Growth Restricted Wistar Rats.

Introduction: Nephrogenesis is the process that leads to the formation of nephrons and ceases around the 36th week of gestation in man, without the possibility of additional formation later in life. A lower number of nephrons have been associated with an increased chance at chronic kidney disease development. In the Netherlands alone, almost 8% of all children are born preterm, and many are treated with drugs or suffer from extra uterine growth restriction (EUGR) that may potentially reduce neonate formation. In this study we investigated the impact of gentamicin and cefazidime on kidney development w/wo EUGR.

Methods: Wistar rats were allocated to either normal size litters (12 pups) or increased size litters (20 pups), the last resulting in EUGR. Both cohorts were divided in control and intervention groups where animals were administered 0.9% NaCl, 4 mg/kg gentamicin or 5 mg/kg cefazidime via intraperitoneal route from post natal day 2-8. At day 8 and 35, animals were sacrificed and the kidneys were collected. Day 8 kidneys were examined for mRNA expression in a selection of targets, proliferation, apoptosis and general histopathology. Total glomerular count (estimated using stereology) and glomerular generation count were performed in kidneys collected at day 35.

Results: Gentamicin treatment in combination with EUGR resulted in 20% less glomeruli compared to sham treatment. EUGR animals had less body weight, but showed parallel growth compared to non-Growth restricted animals, indicating a successful working model. No clear distinctions were noted in mRNA expression levels, glomerular generation count or general histopathology. The proliferation/apoptosis balance is currently under investigation. Conclusion: Early postnatal gentamicin treatment in combination with EUGR tends to decrease the total glomerular number in Wistar rats, of which the pathways were not clarified yet. The long term clinical sequelae are still unclear.
2203 Resistance to Dioxin-Induced Hydronephrosis in a Mouse Strain Having Unresponsive Microsomal Prostaglandin E Synthase-1.

K. Aida-Yasuoka, W. Yoshioka, T. Kawaguchi, S. Ohnoko and C. Tobiyama, University of Tokyo, Tokyo, Japan.

Background: The majority of dioxin toxicity is governed by aryl hydrocarbon receptor (AhR), and the degree of toxicity is affected by its allele type. It is known that AhRb1 and AhRb2 are responsible for marked manifestation of dioxin toxicity, but that AhRd is not. Our previous works 1, 2 showed that COX-2 and microsomal prostaglandin E synthase-1 (mPGES-1), an inducible form of PG2 synthase, are critical factors in the pathogenesis of hydronephrosis (HN), and suggested the presence of the possible strain difference in the development of HN between C57BL/6J and BALB/c. Thus, we here examined the incidence of dioxin-induced hydronephrosis (HN) in mouse pups of these strains to clarify causative factors that bring out a strain difference in dioxin-induced HN.

Methods: Two strains of mice, C57BL/6J and BALB/c, harboring b1 and b2 alleles, respectively, were administered 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD), at an oral dose of 15 μg/kg, 1 day after birth, to expose pups with TCDD via lactation. Kidneys were collected on postnatal day 7 for histology and gene expression analysis by quantitative RT-PCR.

Results: Incidence of HN was approx. 60% in C57BL/6J and 0% in BALB/c despite the comparable induction levels of CYP1A1 in both strains. There was no strain difference in COX-2 mRNA abundance. A strain difference in mPGES-1 mRNA abundance was found comparable to the HN incidence. Gene expression of early growth response 1 (Egr-1), that activates mPGES-1 transcription, was increased in C57BL/6J, but not in BALB/c. In addition, mRNA of aquaporin 2, a water channel, that absorbs water at the collecting duct, was decreased in C57BL/6J and increased in BALB/c.

Conclusions: Although both C57BL/6 and BALB/c have dioxin-sensitive receptors, this study demonstrates that strain difference in the incidence of TCDD-induced HN in mouse pups can be partly explained by distinct expression differences in mPGES-1. It is also suggested that Egr-1 may modulate the induction of mPGES-1.


2204 Escape from Toxic Island: Learning Toxicology Concepts through Informational Posters and a Board Game.

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Escape from Toxic Island is a program that was developed by the MASOT Education and Outreach Committee to teach toxicology concepts to school aged children. The program is composed of informational posters containing basic toxicology concepts followed by a board game with toxicology questions. Answers to many of the questions were contained in the posters and children were encouraged to use them to obtain correct answers as they moved along the game board. One of the informational posters contained basic definitions and toxicology concepts such as routes of exposure, signal words used in the labeling of toxic substances and government agencies that regulate toxicity. The second poster explored toxicity in and around the home and included information about cleaning products, pesticides, and car care products. The program was developed by the Chair of the MASOT Education and Outreach Committee. Review of the informational posters and game board was accomplished by Education and Outreach Committee members who represent academia, industry and government experts in toxicology to ensure relevancy and accuracy of information. In step with the success of the Inspector Tox program developed 2 years ago by this committee, island themed costumes and props were utilized. Toxic Island’s resident pirate, Captain Tox, assisted with instruction and game play. Incorrect answers were cheerfully corrected after “walking the plank”. The audience for the program resulted from a joint effort of St. John’s University and the Afterschool All Stars Program. The Afterschool All Stars program provides free programs to roughly 80,000 children in need in 13 different cities in the continental U.S. and Hawaii. Volunteers were recruited from MASOT, St. John’s faculty, and graduate and undergraduate toxicology students. Learning outcomes for the program are based on responses made to questions posed on the game board. Respondents who answered incorrectly to game board questions were encouraged to explore the informational posters for correct answers.

2205 A C. elegans Dose-Response Protocol and Inquiry Lab in an Undergraduate Toxicology Course.

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In order to prepare students for a multi-week research project, it is essential to allow them to practice the protocols that will be used in the project. Using C. elegans, students initially observed normal behavior and identified the developmental stages of the organisms. Students then practiced transferring C. elegans between plates and explored various endpoints that could be used in their research project, including avoidance, locomotion and feeding. During the next lab session pairs of students were given protocols for a dose-response lab activity in which they were assigned a concentration, performed a dilution series, and used the dry drop test as a measure of avoidance. Data were compiled and statistically analyzed after the lab. After establishing this basis of knowledge the students designed and conducted their own experiment for the remainder of the semester. Student pairs wrote a research proposal based on a literature search, gathered preliminary data in the lab and embarked on several replications of their experiment. Students analyzed their data using an ANOVA and presented their research as a poster session to the broader community.

2206 The Best of the Worst: A Novel Approach to Teaching Environmental Toxicology.

C. P. Curran, Biological Sciences, Northern Kentucky University, Highland Heights, KY.

Curriculum development for undergraduate toxicology courses taught is challenging, because of the varied preparation and background of students enrolled. Students are typically majoring in environmental science, chemistry, or biology and each discipline has unique program requirements. Environmental toxicology courses present even greater challenges for students without a solid grounding in ecology or biochemistry. A novel course was developed at Northern Kentucky University, focused on problem-based learning and team-based learning to engage students in identifying and understanding environmental toxicology issues in their local communities and in a variety of ecosystems (polar, temperate and tropical). Sample activities such as “What is the Worst Environmental Problem in the World?” will be explained along with the pedagogical underpinnings of the curriculum design. Student satisfaction with the course was extremely high (4.8 on a 5.0 scale), indicating this course could be a model for other undergraduate educators.

2207 Risk Assessment Capstone Project for Seniors in an Undergraduate Toxicology Program.

S. M. Ford, College of Pharmacy & Allied Health Professions, St. Johns University, Jamaica, NY.

Capstone projects are intensive, active learning exercises for seniors to apply their knowledge and skills to a complex problem in their discipline. The projects vary in form and function. They may be done in teams or by individual students; the tasks may be self-selected or assigned. Planning and implementation are student-directed under supervision of faculty. Outcome of the work may be a written document and/or an oral presentation. The scope should be substantial, utilize critical thinking, and draw upon the learning objectives of the major. In our BS Toxicology program, seniors are assigned a risk assessment in the course Regulatory Toxicology and Risk Assessment. They are given a hypothetical disaster, involving a population exposed to a chemical through air, water, food, soil, or medication. Ideally the chemical chosen is one for which the toxicological data is sparse, so that students must evaluate the agent based on its properties and chemical class. The scenario includes the amount released, the exposed population, and the media of exposure. The result of the project is a written risk analysis, a website, and presentation of their findings to the College. The capstone project requires the students to utilize the facts and concepts of toxicology in an analytical manner, and apply the skills of writing, oral communication, and teamwork to a realistic situation. The public presentation informs the larger university community on the process of toxic risk assessment.
Development of a Summer Undergraduate Research Program in Toxicology and Environmental Health Sciences.

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Exposure to research opportunities in toxicology and environmental sciences is key to the development of the next generation of scientists. The Community Outreach and Engagement Core of the NIEHS Center for Environmental Exposures and Disease at the University of Medicine and Dentistry of New Jersey-Robert Wood Johnson Medical School and Rutgers University has developed a summer research fellowship program to promote toxicology and environmental sciences as careers in biomedical research. The program consists of 10-week basic science and translational research experiences for undergraduates and was also designed to include weekly events including laboratory safety and responsible conduct of research training, a field trip to a pharmaceutical company, career development and research seminars and student presentations. Participants of the 2012 summer research program ranked the field trip as the most valuable weekly activity followed by presentations from toxicologists and environmental health scientists. Based on pre- and post-survey results, over 60% of respondents reported that a career as a scientific researcher was most appealing based upon satisfaction from doing research, the perceived benefit of scientific knowledge to the community, and an overall interest in science. In addition, 87.5% of respondents will continue to pursue research after completion of the summer research program. This includes five students pursuing Ph.D. degrees beginning in 2012 or 2013. A summer research program engages undergraduate students in full-time research experiences and provides unique opportunities to promote toxicology and environmental sciences as research areas for the next generation of scientists and enhances career development skills. Supported by ES020721, ES050022, and ASPET SURF.

Incorporation of Toxicology and Risk Assessment Principles into an Environmental Health Course.


Environmental health deals with a multitude of public health issues ranging from sanitation to hazardous materials to air and water quality. Practitioners are largely involved in the regulatory side of protecting public health, which requires an intimate understanding of toxicology and risk assessment. However, many of the identified deficiencies with risk assessment are perpetuated by the educational curricula. For example, chemical hazards are introduced to students and evaluated by regulators on a case-by-case basis, often based on one toxicological endpoint alone. Strong emphasis is placed on making comparisons of reference doses to exposure estimates without regard to the effect of uncertainties associated with inadequate data. High uncertainty alone could result in selection of a more toxic substance over one less studied. A unique class project was created to give environmental health students, our future regulators, an improved perspective on how toxicological information can be used to make informed decisions that best protect human health and well-being. The multi-step exercise focuses on a decision made by the World Health Organization to bring DDT back into use to fight malaria. Students work in groups to collect chemical, toxicological, efficacy, and economic aspects of a common pesticide used for mosquito control. The class then comes together to compare and discuss the group results and select the best pesticide option for the impacted region. A semi-quantitative risk management tool is also used for a more objective determination. Lastly, a surprise change in one circumstance causes perspectives to change and the outcome is a completely different conclusion. The end result is a sophisticated interaction of ideas and concepts, which motivates students to start thinking about how much is more to toxicology than lethal doses and dose-response curves, and that there is more to risk assessment than hazard quotients. A detailed description of the project and handouts will be provided.

Poisoning Principles: Clinical Toxicology and Undergraduate Nursing Education.

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According to the American Association of Poison Control Center’s (AAPCC) annual report (2010), 601,197 human exposures to toxicants resulted in management in a health care facility. Therefore, education concerning clinical toxicology and poisonings is essential in preparation of future healthcare professionals. This abstract describes a program for educational enrichment for undergraduate nursing students through a collaboration with a regional Poison Control Center (PCC) and a local university. Undergraduate nursing students were assigned to a clinical toxicology rotation and were required to participate in didactic educational modules addressing principles of toxicokinetics, such as the ADME (absorption, distribution, metabolism, and exposure) model, prior to the PCC rotation. During the rotation, clinical toxicology approaches to the poisoned patient were reviewed. Attendance at a journal club at the poison center was included, covering topics such as: environmental exposures, chemicals, drug overdoses, and unintentional poisonings. A lecture on lead toxicity was then given to the entire class, including those undergraduate students in other rotations. The program culminated with all students participating in a poster session addressing clinical toxicology and other environmental health experiences. Students were required to evaluate the entire course, including the PCC rotation. Students reported an increase in knowledge and awareness about the role of toxicology and environmental health in their professional development. In addition, the knowledge gained has heightened the role of poisoning and antitox use, along with toxicology principles, for treatment and care of patients utilizing healthcare facilities. Recommendations for future programs include integrating toxicology into didactic and clinical experiences of under-graduate nursing students within all rotations, and across healthcare and scientific disciplines.


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Over 42% of the estimated 130 million tons of coal ash produced annually in the US is beneficially used, which is material that is not placed in disposal facilities. However, these uses are threatened by USEPA’s potential regulation of coal ash as a hazardous waste, and by constant references to “toxic coal ash” by the press in response to environmental groups’ writings. Therefore, a detailed health-risk based evaluation was conducted of coal ash data released in a report by the US Geological Survey. Eight robust coal ash data sets were selected as representing material that could be put into beneficial use from five US power plants each utilizing a different source of coal. The evaluation was conducted by comparing constituent concentrations in coal ash to risk-based screening levels developed by the USEPA that are protective of a child’s direct exposure to residential soils (including ingestion, dermal contact and inhalation routes of exposure). These screening levels are considered by the Agency to be protective for daily exposure by humans (including sensitive groups) over a lifetime. Coal ash percentiles (10th-90th) were compared directly to the screening levels, and in a more detailed evaluation, upper-bound exposure point concentrations were used in a cumulative risk screening process. Constituent concentrations in coal ash were also compared to USGS background concentrations in soils in the US. The results indicate that with few exceptions constituent concentrations in coal ash are below screening levels for residential soils, and are similar in concentration to background US soils. Because exposure to constituents in coal ash used in beneficial applications, such as concrete, road base, or structural fill would be much lower than assumed for a residential soil scenario, these uses should also not pose a direct contact risk to human health.

Provisional Advisory Levels (PALs) for Chloroethanol (Ethylene Chlorohydrin).

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PAL values developed for hazardous materials by the US EPA represent general public emergency exposure limits for oral and inhalation exposures corresponding to three different severity levels (1, 2, and 3) for 24-hour, 30-day, 90-day, and 2-year durations. PAL 1 represents the threshold for mild effects; PAL 2 represents the threshold for serious, irreversible or escape-imparing effects; PAL 3 represents the threshold for lethal effects. PALs have not been promulgated nor have they been formally issued as regulatory guidance, but are intended for use at discretion of risk managers in emergency situations when site-specific risk assessments are not available. The PAL document for chloroethanol was developed based on the SOP and QAPP requirements. Chloroethanol (2-chloroethanol; ethylene chlorohydrin), is a colorless, combustible liquid with a faint ether-like odor, and is soluble in water and organic solvents. It is a high production volume chemical that is used as an industrial solvent and a chemical manufacturing intermediate. It is formed when ethylene oxide is used to sterilize polyvinyl chloride plastics and to fumigate foods. Humans and animals exposed by inhalation or orally exhibited CNS, GI, and respiratory symptoms, and had post-mortem lesions in numerous internal organs. All PALs were derived using rat
data because the human data were not adequate; the animal data were generally consistent among species and with the limited human data. Developed PAL values for oral exposure are 35 mg/L as the PAL 1 for 24 hours; 110 mg/mL as the PAL 2 for 24 hours, and 30 and 90 days; 230 mg/L as the PAL 3 for 24 hours; and 160 mg/L as the PAL 3 for 30 and 90 days. PALs derived for inhalation exposure are 0.26 ppm and 0.78 ppm as the 24-hour PAL 2 and PAL 3, respectively. For 30 and 90 days, the inhalation PAL 1 is 0.0015 ppm and the PAL 2 is 0.015 ppm. Other oral and inhalation PAL values were not developed due to insufficient data.

2213 Provisional Advisory Level (PAL) Development for Lewiste and Sulfur Mustard.


PAL values developed for hazardous materials by the US EPA represent general public emergency exposure limits for oral and inhalation exposures corresponding to three different severity levels (1, 2, and 3) for 24-hr, 30-d, 90-d, and 2-yr durations. PAL 1 represents the threshold for mild effects; PAL 2 represents the threshold for serious, irreversible or escape-impairing effects; PAL 3 represents the threshold for lethal effects. PALs have not been promulgated nor have they been formally issued as regulatory guidance. They are intended to be used at the discretion of risk managers in emergency situations when site specific risk assessments are not available. Application of PAL protocols has been performed for lewiste and sulfur mustard to estimate inhalation exposure limits; oral PAL values are not recommended (NR) due to insufficient data. PAL values for the vesicants, lewiste and sulfur mustard, are based on ocular effects in humans and animals or lethality thresholds in rodents.

Lewiste inhalation PAL 1, 2, and 3 values are NR, 0.01, and 0.037 mg/m3, respectively, for 24-hr; and NR for 30-90-d and 2-yr.

Sulfur mustard inhalation PAL 1 values are 0.00083 mg/m3 for 24-h and 0.00010 mg/m3 for 30-90-d and 2-yr. PAL 2 values are 0.0042 mg/m3 for 24-h, 0.0009 mg/m3 for 30-d, and 0.00097 mg/m3 for 90-d and 2-yr. PAL 3 values are 0.088 mg/m3 for 24-h and NR for 30-90-d and 2-yr.

2214 Provisional Advisory Level (PAL) Development for Fentanyl.

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PAL values developed for hazardous materials by the US EPA represent general public emergency exposure limits for oral and inhalation exposures corresponding to three different severity levels (1, 2, and 3) for 24-hr, 30-d, 90-d, and 2-yr durations. PAL 1 represents the threshold for mild effects; PAL 2 represents the threshold for serious, irreversible or escape-impairing effects; PAL 3 represents the threshold for lethal effects. Minimum data requirements must be met or a value may be considered NR (not recommended). PALs have not been promulgated nor have they been formally issued as regulatory guidance. They are intended to be used at the discretion of risk managers in emergency situations when site specific risk assessments are not available. Application of PAL protocols has been performed for fentanyl to estimate oral and inhalation exposure limits.

Fentanyl is a highly potent, synthetic opioid used clinically as an analgesic and anesthesia. It is currently analyzed using GC-MS. Partially metabolized fentanyl to estimate oral and inhalation exposure limits.

2215 Development of a Chronic Reference Concentration for Decalin.

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Decalin is a naturally occurring dicycloalkane present in crude oil and produced as a product of combustion. Its ability to solubilize oils and fats makes it useful in paints, cleaning fluids, gasoline, and varnishes. Because of its widespread use in many types of commercial products, decalin is ubiquitous in the environment and exposure by the general public is of concern. Neither oral nor inhalation toxicity values are currently available for decalin in published sources despite recurring point and non-point source releases. To derive a reference concentration (RfC) for decalin, inhalation toxicity studies were reviewed using a weight-of-evidence approach. A two-year mouse inhalation study conducted by the National Toxicology Program was chosen as the critical study for the derivation of the chronic RfC. The results of this study, a significant increase in the occurrence of hepatotoxicity was detected at the highest concentration tested (400 ppm) in male mice. Benchmark dose modeling was utilized to derive a point of departure for hepatic necrosis, syncytial alteration, and erythropagocytosis. For data not amenable to modeling, a point of departure was derived using the no observable adverse effect level (NOAEL). The most sensitive adverse effect, syncytial alteration resulted in a BMDL10 of 7.8 ppm using the Log-logistic model. A chronic RfC for decalin of 0.08 mg/m3 was calculated by conversion of the BMDL10 to a human equivalent continuous inhalation concentration of 1.4 ppm (7.9 mg/m3) using a dosimetric adjustment factor of 1 and application of a total uncertainty factor of 100. The chronic decalin RfC was derived despite several toxicity database limitations, including a small number of chronic inhalation studies and uncertainty regarding reproductive effects. Future research on decalin toxicity is needed to better characterize the adverse effects associated with its chronic inhalation.

2216 Weight-of-Evidence Evaluation of Methyl Methacrylate Olfactory Effects in Humans and Derivation of an Occupational Exposure Level.

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Methyl methacrylate (MMA) causes olfactory effects in rodents that are considered relevant to humans. Recent scientific studies have focused on understanding the apparent lack of species concordance between the rodent and occupational studies. We have applied the hypothesis-based weight-of-evidence (HBWoE) approach to evaluate the concordance of the available data and the hypothesis that the observed difference in sensitivity between rats and humans may be the result of physiological and biochemical differences. Our WoE analysis integrated several lines of evidence (animal, human, mode of action (MoA), and toxicokinetics data) and found: 1) acute and chronic rat and mouse MMA inhalation studies consistently indicate degenerative lesions of the main olfactory region as the most sensitive endpoint; 2) numerous studies support an MoA for MMA involving high concentrations of carboxylesterase activity in nasal epithelial tissue that metabolizes MMA to methyl acrylic acid (MAA), an organic acid with irritative and corrosive properties; 3) carboxylesterases are a group of non-specific enzymes that are widely distributed in the human and animal olfactory systems; 4) toxicokinetic studies and a physiologically based pharmacokinetic (PBPK) model describing inhalation dosimetry throughout the body in animals and humans; 5) toxicokinetic studies and a physiologically based pharmacokinetic (PBPK) model describing inhalation dosimetry throughout the body in animals and humans; 6) occupational exposure levels (OELs) from animal data (ranging from 28-118 ppm) and human data. Overall, our WoE analysis supports use of the human data for derivation of an MMA OEL of 50 ppm.

2217 An Updated Dose-Response Evaluation of Aldrin and Dieldrin.


The United States Environmental Protection Agency (USEPA) reviewed the cancer and non-cancer effects of the organochlorine insecticides, aldrin and dieldrin, in the late 1980’s. The results of these assessments are reported by USEPA’s Integrated Risk Information System (IRIS). These assessments include reference dose (RfD) values of 0.00003 and 0.00005 mg/kg-day for cancer slope factor values of 17 and 16 (mg/kg-day)-1 for aldrin and dieldrin, respectively. The USEPA methods for dose-response analysis have changed in the decades since these evaluations were done. The dose-response analyses of cancer and non-cancer health effects of aldrin and dieldrin were re-evaluated using current methodology, including benchmark dose (BMD) analysis (BMDS Version 2.2 software) and current body weight scaling. A literature review was updated to determine the most appropriate adverse effect endpoints. Using current methodology and information, the cancer slope factors for aldrin and dieldrin were estimated to be 3.4 and 7.0 (mg/kg-day)-1, respectively (i.e., about 5 and 2.3 lower risk than previous assessments). The current analyses estimated RfD values of 0.0001 and 0.00008 mg/kg-day for aldrin.
and dieldrin, respectively (both higher than previous assessments). Because aldrin and dieldrin are no longer used as pesticides in the United States, they are a low priority for additional review by the USEPA. However, because they are persistent and still detected in environmental samples, quantitative risk assessments based on the best available methods are required. Several national and international health assessment organizations (e.g., WHO) do not consider aldrin and dieldrin to be human carcinogens. Recent epidemiologic studies do not demonstrate a causal association between aldrin and dieldrin and human cancer risk. These re-evaluations, based on current methodologies and available data, suggest that these two compounds pose a lower human health risk than currently reported by USEPA.

2218 Benzene: Development of a 24-Hour, Health-Protective Comparison Value.
J. T. Haney, Toxicology Division, Texas Commission on Environmental Quality, Austin, TX.

Texas has the most extensive volatile organic compound (VOC) ambient air monitoring network in the nation. As part of that network, the TCEQ collects every sixth-day, 24-hour canister VOC data. These data are used to calculate annual averages for comparison to chronic, health-protective Air Monitoring Comparison Values (AMCVs) (i.e., RfC-like values for noncarcinogenic effects, 1E-05 excess risk levels for cancer effects). In regard to acute exposure durations, however, the TCEQ typically has only 1-hour AMCVs, which while conservative are not designed to evaluate 24-hour sample results. Thus, the development of 24-hour, health-protective AMCVs would allow the TCEQ to more fully utilize 24-hour VOC data for the evaluation of potential public health concerns. The TCEQ has developed a proposed 24-hour AMCV for benzene since it is a ubiquitous VOC of both agency and public interest. Critical effect dose-response data for hematotoxicity from mouse studies indicate an effect level range of 10-100 ppm for subacute exposure (e.g., 6-8 hours per day, 5-10 days). A point of departure (POD) from these studies was used to develop the 24-hour value. The total number of exposure hours exceeds 24 hours for all these subacute studies and available toxicokinetic information indicates the time between the intermittent daily exposures would not allow for clearance of benzene's hematotoxicity-implicated metabolites (e.g., hydroquinone, hydroquinone glucuronide, catechol) from the bone marrow as evidence suggests they are not readily excreted. Using the same POD (LOAEL of 10.2 ppm) and an uncertainty factor of 100 (UF of 1000) for the TCEQ used to determine the 1-hour AMCV (180 ppb) but without duration adjustment (based on toxicokinetic considerations) results in a conservative 24-hour, health-protective AMCV of 100 ppb. This value is well below even chronic human hematotoxicity observed adverse effect levels (e.g., 7.2-13.6 ppm). The proposed 24-hour AMCV is considered sufficiently conservative for the adequate protection of public health and would significantly complement TCEQ health effect evaluations of ambient air data.

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It has been demonstrated that swallowed jewellery items may become lodged in the stomach, and morbidity and mortality have been associated with jewellery containing lead. Due to its inherent toxicity, the use of cadmium to make costume jewellery may pose an analogous threat. In the absence of reliable human toxicity data, the results from animal studies were used to derive an oral acute provisional minimal risk level (pMRL) of 0.0732 mg/kg bw for cadmium. Health Canada analyzed approximately 200 children’s jewellery samples that were judged small enough to fit into a child’s mouth for cadmium content. A subset of these samples were also subjected to migration testing, which revealed no consistency between the total amount of cadmium in a sample, and the amount that might be released in the simulated physiological environment of the stomach over an extended period (such as in the case of a piece of jewellery lodged in the stomach over several days). Since standardized migration testing cannot accurately predict the amount of cadmium that might leach out of a sample under such circumstances, the use of total cadmium content to derive a guideline is considered the most health-protective approach. Limiting total cadmium content to 130 ppm removes variables that interfere with migratable cadmium quantification (such as the thickness and composition of the surface plating, and elemental composition of the amalgam) and is protective of the scenario of an ingested piece of jewellery lodged in the stomach for an extended period of time.

2220 Assessment of Risks to the US Population Posed by Exposure to Gold and Ceramic Dental Restorations.
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There is significant mercury exposure from dental amalgam used in restorative practice. However, little is known about the chemical exposures and risks of alternative dental restorative materials. Thus, it is difficult for clinicians to weigh the performable, risks, and benefits of dental amalgam to alternate restorative materials. Here we provide the first population-level risk assessment for gold alloy and ceramic restorative materials. Employing the US National Health and Nutrition Examination Survey (NHANES) data from 2001 to 2004, we assessed the exposure of adults to the components of gold alloy and ceramic dental restorations in the US general population. Three specific exposure scenarios were considered: 1) all restorations were either gold alloy or ceramic; 2) all crowns were gold alloy or ceramic; and 3) 11% of fillings were either gold alloy or ceramic, in 30% of the population. Silver appears to be the most problematic component of gold alloy restorations, due to a combination of relatively high toxicity and high proportional composition. Based on the toxicity of silver and its proportional content in gold dental alloys, it was estimated that adults could possess an average of 16 tooth surfaces restored with gold before exceeding the Reference Exposure Level (REL) for silver. Lithium appears to be the most problematic component of dental ceramics. All other ceramic components considered resulted in estimated daily doses well below their respective RELs. Based on the toxicity of lithium and its proportional content in dental ceramics, it was estimated that adults could possess an average of 4 tooth surfaces restored with ceramics before exceeding the REL for lithium. Relative to dental amalgam and gold alloys, ceramics present the fewest and lowest chemical exposures and risks.

2221 Development of an Oral Cancer Slope Factor for Acrylamide Based on Tumors Relevant to Humans.
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Acrylamide is an industrial chemical used mainly in the production of polycrylamides. Acrylamide and polycrylamides have many uses, including uses as flocculants and flow control agent for enhancing oil production from wells, in the production of dyes, organic chemicals, contact lenses, cosmetics and toiletries, in sugar refining, and as a chemical grouting agent and soil stabilizer. Acrylamide is also commonly used in fried foods and the potential routes of human exposure in the general population is diet. In their most recent risk assessment for acrylamide, USEPA developed an oral cancer slope factor (OSF) of 0.5 (mg/kg-day)-1 based on a 2-year drinking water study that reported increased incidences of thyroid tumors and tunica vaginalis mesotheliomas (TVMs) in male F344 rats. However, there is considerable evidence that F344 rats are particularly susceptible to TVMs, and therefore TVMs may not be relevant to humans. As such, our objective was to evaluate the overall weight of the evidence regarding each tumor type and to derive an OSF for acrylamide based on those tumors relevant to humans. Among the tumors induced by acrylamide (thyroid tumors, TVMs, mammary gland tumors, CNS tumors), thyroid and mammary gland tumors were considered most relevant to humans. Using the rat OSF for the combined increased incidences of thyroid and mammary gland tumors observed in female F344 rats, we derived a human OSF using a rat-to-human dose metric conversion factor based on serum levels of the primary acrylamide metabolite, glycidamide (widely considered to be the putative proximal carcinogen). The final human OSF for combined thyroid and mammary gland tumors was determined to be 0.09 (mg/kg-day)-1. This OSF suggests a lower cancer potency for acrylamide based on target tissues more relevant to humans. It should be noted however, the FDA and NTP just completed a 2-year, multi-species drinking bioassay and the results of this study may impact future OSF estimates for acrylamide.

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The carcinogenic potency of MTBE was initially evaluated by State of California scientists in 1999 based on a gavage study with rats (Bellego et al. 1995, 1997, 1998) and inhalation studies with rats and mice (Bird et al. 1997). Potency was estimated at that time using a linearized multistage model. Since then, an additional
rodent cancer bioassay of MTBE in drinking water (Dodd et al. 2011), several genotoxicity and mutagenicity studies, evaluations of MTBE metabolites formaldehyde and tert-butyl alcohol (TBA), as well as other studies providing information on MTBE’s mode of action have been reported. In addition, the U.S. Environmental Protection Agency (EPA) declared the data on lymphomas and leukemias from Belpoggi et al. (Ramazzini Institute) as unreliable for risk assessment raising additional uncertainty about California’s potency estimate. The new data and remaining reliable historic rodent bioassay data were used to re-evaluate the cancer potency or slope factor (CSF) of MTBE considering mode of action (MOA). The overwhelming majority of studies indicate that neither MTBE nor TBA is genotoxic. In stark contrast, formaldehyde is clearly genotoxic, but when generated by MTBE metabolism is efficiently detoxified to preclude elevating background levels and associated genotoxic damage. Based on genotoxicity and MOA analyses, the most likely CSF for MTBE is zero, unless chronic exposures induce target-tissue toxicity including in sensitive individuals. A corresponding expected CSF value for MTBE conditional on a linear MOA was estimated to be 0.000018 mg MTBE per kg body weight per day for a chronically exposed adult. If MTBE is carcinogenic to humans, then it is extremely weak relative to other chemical carcinogens evaluated by EPA.

2223 Development of a 24-Hour Air Monitoring Comparison Value (AMCV) for 1, 3-Butadiene and Comparison of 1-Hour, 24-Hour, and Chronic AMCVs to Observed Adverse Effect Levels.

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The TCEQ develops AMCVs, which are considered safe concentrations of chemicals in air, to determine whether 1-hour (hr) or annual average chemical concentrations in ambient air exceed levels of potential concern for adverse health effects. Previously for 1,3-butadiene (BD), a 1-hr AMCV of 1,700 ppb was derived based on a mouse developmental study (6 hr/day exposures, gestational days 6-15). A chronic AMCV of 9.1 ppb was derived based on a 1 in 100,000 excess risk for leukemia mortality from an epidemiological study in styrene-butadiene workers. To calculate annual ambient air concentrations, the TCEQ collects a significant amount of 24-hr monitoring data that are not directly comparable to the 1-hr or chronic AMCV. Therefore, the TCEQ has developed a 24-hr AMCV for BD to evaluate the potential for health effects from 24-hr exposure. The same mouse study used for the 1-hr AMCV was judged to be the critical study for the 24-hr AMCV based on mode-of-action and toxicokinetic data. The 6-hr human equivalent point of departure (POD-HEC) of 51,300 ppb was duration adjusted (Haber’s rule with n = 1) to calculate the 24-hr POD-HEC of 12,800 ppb. The proposed 24-hr AMCV is 430 ppb after application of total uncertainty factors of 30. The TCEQ has developed guidelines to derive observed adverse effects levels (OAELs) based on available dose response data to better communicate to risk managers and the public the margin of safety between OAELs and AMCVs. For the 6- and 24-hr AMCVs, the acute OAEL is 66,000 ppb (the central estimate POD-HEC for 5% decrease in fetal weight loss). The 1-hr AMCV of 1700 ppb is 39 times lower than the acute OAEL and the 24-hr AMCV of 430 ppb is 150 times lower. The long-term OAEL is 10,000 ppb (the lowest average occupational exposure concentration where the likelihood ratio test that slope = 0 was statistically significant for different maximum levels of cumulative BD ppm-years for leukemia deaths). The chronic AMCV of 9.1 ppb is 1,100 times lower than the chronic OAEL.

2224 Health-Based 24-Hour Air Monitoring Comparison Value (AMCV) for Acrrolein in Ambient Air: Comparison to Air Monitoring Data Collected in Texas.

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Acrrolein is of national and state interest because it is ubiquitous, is difficult to analyze in ambient air, and concentrations causing eye and respiratory irritation are low. In 2011, a follow-up special monitoring project for acrolein was conducted by the USEPA at a school near a building products manufacturing facility in Texas. Ten 24-hour (hr) ambient air samples were collected in canisters downwind of a facility known to emit acrolein and analyzed by GC/MS to better evaluate the potential for adverse health effects. Acrrolein’s toxicity is mainly concentration dependent and levels causing adverse effects are similar in humans and animals. The same rat study used to develop the chronic AMCV was selected as the critical study for derivation of the 24-hr AMCV since 24-hr exposure levels were permissive of exposure durations (e.g., 6 hr/day for 4, 14, 30, and 65 days) which encompassed the 24-hr duration of interest. Therefore, the 24-hr AMCV was derived based on the 1 in 100,000 excess risk for leukemia deaths. A no-observed-adverse-effect level of 200 ppb was identified from the key study for all exposure durations based on the absence of nasal epithelial hyperplasia. Based on a mode-of-action analysis, no duration adjustment was necessary. After correcting for animal-to-human dosimetric differences, the 24-hr human equivalent point of departure was 37.4 ppb. Total uncertainty factors of 30 were applied to calculate the 24-hr AMCV of 1.2 ppb. In comparison, the 1-hr AMCV for acrolein is 4.8 ppb and its chronic AMCV is 0.22 ppb. No concentrations exceeding the 24-hr AMCV using the improved canister method have been reported at monitoring sites in Texas.

Millions of people worldwide are exposed to arsenic (As) via drinking water. As is biotransformed for excretion and the CH3 groups necessary for such metabolism are also essential to DNA methylation. Pregnant women methylate As more efficiently to protect the fetus from immediate toxicity, but at the risk of latter epigenetic deregulation. Thus, preventing early-life As exposure may be critical for the health of future generations. In order to evaluate early-life exposure to arsenic, pregnant hamsters were exposed to 50 and 100 ppm of arsenite via drinking water at the 0th and 9th day of gestation. Co-treatment with 8.5 mg/Kg/day of selenium or 6 mg/Kg/d of vitamin E was also carried out to assess whether those compounds exert any As-transplacental protective role. To characterize arsenic speciation during pregnancy, fetus development, unwrapping and adulthood, As species were measured in the urine and or organs by HPLC/ICP/MS. Teratogenic effects of transplacental arsenic were analyzed observing both reabsorptions and embryo anomalies, and in utero As-exposed litters were followed-up to evaluate tumor incidence at the adult stage. Characterization of arsenic species indicates that fetuses and unwrapped offspring are As-exposed through placenta and milk, showing a speciation profile similar to that in mothers. Results demonstrate a teratogenic effect as a dose and time-of-gestation dependent manner, and a noticeable protective effect of both selenium and vitamin E.

To provide perspective on current and proposed occupational exposure limits (OELs) for peracetic acid (PAA; CAS 79-21-0) we evaluated PAA toxicity with the aim of understanding uncertainties and their implications for the resulting OEL. The database for PAA is limited and no single study is definitive. Two unpublished reports on human exposures to PAA provide some concentration-response data, indicating that a sensitive acute effect of PAA exposure is eye and respiratory tract irritation, but the studies differ quantitatively. These differences are not surprising, in light of the differences in exposures (apparently pure PAA vapor vs. an aerosol of a mixture), the subjective nature of the reporting, and the likely small sample sizes. The studies are also limited by the lack of clear concentration-duration-response data. Nonetheless, the studies provide a reasonable estimate of the threshold for the onset of irritation in humans in the range of 0.53 mg/m3 for up to 3 hours and 1.56 mg/m3 for up to 45 minutes. RD50 (concentration estimated to cause a 50% depression in respiratory rate) data in mice and rats provide additional information on the irritant potency of PAA. The RD50 in mice was 17 mg/m3 for pure PAA vapor and 12 mg/m3 for a commercial mixture. The rat RD50 was 21.5 to 24.1 mg/m3. Based on the array of human data and the RD50 values in rodents, we calculated potential TWA OELs ranging from 0.26 to 1.56 mg/m3. A similar ratio of 0.62 – 2 mg/m3 is found among the published OELs, and any of these values could be justified as protective of worker health given the uncertainties in the data and the precision of the OEL methodology. More definitive sensory irritation studies would further clarify selection of a value in this range. Given the extant data, the ultimate OEL, choice within a range of reasonable values is a policy-based risk management decision that cannot be based on science alone. The current time averaging approach is also not clearly established by the data; however, a combination of a TWA with a STEL is the recommended risk management option.
An Inhalation Risk Assessment for Measured Ambient Air Concentrations of 6:2 Fluorotelomer Alcohol.


6:2 Fluorotelomer Alcohol (CAS #647-42-7, 1-Octanol-3,4,4,5,5,6,7,8,8,8 tridecafluoro-, 6:2 FTOH) is a raw material used for manufacturing surfactant and polymeric products. 6:2 FTOH vapor phase inhalation is a potential exposure route. The aim of the current investigation was to 1) compare the oral and inhalation repeated-exposure toxicity data to confirm systemic toxicity, target organs, and lack of an exposure route effect, 2) confirm similar metabolic and toxicokinetic profiles via both exposure routes, and 3) conduct an inhalation risk assessment for reported ambient air concentrations. In an inhalation range-finder (5-days) and a 28-day inhalation toxicity study, the profile of 6:2 FTOH and its metabolites in plasma under controlled inhalation exposure was investigated as well as the systemic toxicity and target organs. These studies provided a basis for toxicity comparison, plasma metabolites, and dosimetry between inhalation and oral dosing. Similar toxicity, metabolic and toxicokinetic profiles via both exposure routes was confirmed. Benchmark Dose Analysis (BMD) was conducted on the subchronic toxicity endpoints to determine the most sensitive effect and the corresponding BMD associated with this effect. Based on this analysis, the corresponding human equivalent dose (HED) was calculated to be 1.4 mg/kg bw/day. An additional assessment factor of 2 was applied to extrapolate from the subchronic to a chronic exposure and resulted in a final HED of 0.7 mg/kg bw/day. An equivalent air concentration was determined using an allometric scaling factor to arrive at a human equivalent concentration (HEC) of 2.5 mg/m³. This HEC was then divided by the reported indoor and outdoor air concentrations to arrive at a margin of exposure (MOE). MOEs calculated for inhalation exposure to indoor or outdoor air ranged from 1.1E+05 to 2.5E+07. This assessment indicates there is no human health risk expected even at the highest ambient air concentrations of 6:2 FTOH reported.

Derivation of an Occupational Exposure Limit for Inorganic Borates Using a Weight of Evidence Approach.

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Inorganic borates are encountered in many settings worldwide, spurring international efforts to develop exposure guidance (U.S. EPA 2004; WHO 2009; ATSDR 2010) and occupational exposure limits (OEL) (ACGIH 2005, MAK 2011). We derived an updated OEL to reflect new data and current international risk assessment frameworks. We assessed toxicity and epidemiology data on inorganic borates to identify relevant adverse effects. International risk assessment frameworks (IPCS 2005; IPCS 2007) were used to evaluate endpoint candidates: reproductive toxicity, developmental toxicity, and sensory irritation. For each endpoint, a preliminary OEL was derived and adjusted based on consideration of toxicokinetics, toxicodynamics, and other uncertainties. Dose-response modeling supported selection of the point of departure for each endpoint. Developmental toxicity was the most sensitive systemic effect. An OEL of 1.6 mg B/m³ was estimated for this effect based on a point of departure (POD) of 63 mg B/m³ with an uncertainty factor (UF) of 40. Sensory irritation was considered to be the most sensitive effect for the portal of entry. An OEL of 1.4 mg B/m³ was estimated for this effect based on the identified POD and an UF of 1. Reproductive effects are not the most sensitive basis for OEL derivation. An OEL of 1.4 mg B/m³ was derived as an 8-hour TWA based on sensory irritation potential. The OEL is expected to protect from systemic toxicity endpoints.

US EPA Decabromodiphenyl Ether Alternatives Hazard Assessment Results.

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The U.S. Environmental Protection Agency (US EPA) Design for Environment (DfE) Program undertook a chemical alternatives assessment for decabromodiphenyl ether (decaBDE) as part of the Action Plan for Polybrominated Diphenyl Ethers (PBDEs) published in December 2009. DfE convened a multi-stakeholder partnership to explore the human health and environmental profiles of functional and viable alternatives to decabromodiphenyl ether (decaBDE). The partnership identified ~ 30 functional alternatives to decaBDE. The hazard assessment for decaBDE and the alternatives used DfE hazard evaluation DfE hazard assessment frameworks for human health, ecological toxicity and environmental fate endpoints. The alternatives included a range of flame retardant chemicals including both halogenated and non-halogen organic substances, inorganic matter, and polymeric and non-polymeric substances and novel, new to market substances. Some alternatives were well characterized for all endpoints, while others were lacking data. Analog data, predictive models, structural alerts and expert judgment were used to make hazard designations for endpoints with data gaps. Trends for human health, ecological toxicity and fate characteristics were indicated in a matrix of multiple alternatives and endpoints used the same approach. In novel, molecular size ranges, molecular structures, and/or functional groups were found to be most influential on the hazard designations. A novel component of this assessment was the evaluation of higher molecular weight polymers for their human health and ecological toxicity based on their low potential for bioavailability and variability in low molecular weight weight polymers. Effective hazard assessment approaches of hazard criteria coupled with decision-making protocols are practical tools for businesses to use early in materials selection processes and will contribute to more sustainable product development. The resulting hazard profiles should be of value to manufacturers making substitution decisions in preparation for the upcoming decaBDE phase out.

Predicting Bioavailability of Arsenic in Mining Soils.

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Arsenic (As) is a naturally occurring element in soil and a key chemical of concern at former mine sites in California. Risk assessment calculations typically utilize default oral toxicity values, which are based on ingestion of readily soluble forms of As such as sodium arsenate (NaAs). However, mining soils in California are relatively high in iron hydroxide phases that bind As strongly, resulting in reduced solubility/bioavailability. The juvenile swine model is an approved, but often cost prohibitive, method for determining the relative bioavailability (RBA) of As in soils compared to that of NaAs. RBAs can be used to adjust toxicity criteria, resulting in a more accurate site-specific risk assessment. In vitro methodologies have proven to be useful surrogates for in vivo feeding studies in predicting bioavailability for other metals but lack precision for arsenic, particularly in high iron content soils. Six soil samples collected from Empire Mine State Historic Park (total As 302-12,041 mg/kg) were analyzed in the juvenile swine model. RBA's ranged from 4 to 20%. Gastrointestinal model correlated but underestimated RBA (1-9%). Sequential chemical extraction procedures (SEP) were applied to fractionate the As in soils into (F1) non-specifically sorbed; (F2) specifically sorbed; (F3) amorphous and poorly-crystalline oxides of Fe and Al; (F4) well-crystallized oxides of Fe and Al and residual As phases. The results of these extractions demonstrated that the sum of non-specifically sorbed and specifically sorbed arsenic (F1+F2) was similar to the predicted in vitro bioaccessibility while F1+F2+F3 is a conservative estimate of the in vivo RBA (10-50%). SEP could prove to be a cost-effective and valuable screening tool for estimating in vivo RBA. In summary, the assumption of 100% bioavailability of As in mining soils grossly overestimates exposure and risk to human health. Adjustments for As bioavailability in these materials and similar mining wastes provides a more accurate assessment of human exposure.

A Quantitative Risk Assessment of 1-Bromopropane, Based on Tumor Data.

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The “green” movement has resulted in the introduction of several new “environmentally-friendly” substitutes into commerce, including 1-bromopropane (1-BP, CAS no. 96-44-5). Although use of 1-BP is intended to minimize ozone depletion, occupational exposure is of concern. Case studies, occupational exposure assessments, and epidemiological investigations have suggested that workplace exposure to 1-BP may be associated with neurological, reproductive, and hematological effects. Previous quantitative risk assessments of 1-BP have been based on toxicological studies of these and other non-cancer endpoints. This poster presents a quantitative risk assessment based on a NTP chronic bioassay, in which rats and mice were exposed to 125-500 or 62.5-250 ppm 1-BP, respectively, for up to 2 years. Inhalation of 1-BP produced alveolar/bronchiolar adenomas and carcinomas in female mice, adenomas of the large intestine in female rats, and keratoacanthoma/squamous cell carcinoma of the skin in male rats. Benchmark concentrations (BMC) and lower 95% confidence limits (BMCL) estimates at the 1 in 1000 response level (0.1%) were based on a previously published model average procedure. The BMC (BMCL) estimates were 0.85 (0.41 ppm for alveolar/bronchiolar adenoma + carcinoma; 13.5 (2.76) ppm for large intestine adenomas; and 3.73 (1.44) ppm for keratoacanthoma + squamous cell carcinoma of the skin. The BMC
and G.

ily represent the views of the National Institute for Occupational Safety and Health.

lung tumors; 6.17 (1.26) ppm for intestinal tumors; and 1.75 (0.68) ppm for skin

weights, and chronic-active bronchiolar-alveolar inflammation after 13-wks of ex-

mediastinal and tracheobronchial lymph nodes, higher absolute and relative lung

concentration of 8 mg/m3 was estimated to be 0.47 mg/gm lung/day in rats. Based on

accounts for exposure of the general population to bromide from the diet.

traspecies variability, an RfD of 0.7 mg/kg-day was determined for bromide, which

NOAEL was 7 mg Br/kg-day. Using a 10x uncertainty factor to account for in-

sence of sedation and EEG changes within normal limits, the human systemic

were administered NaBr capsules for 12 weeks, there was a small effect on EEGs

cal experience with various bromide salts based on their historical use as sedative-

(unstable) ions, this assessment applies specifically to inorganic bromide, whereas

Dawley rats were exposed 6 h/day, 5 d/wk, for two- or 13-wks to ACUDYNE™

polymer concentrations of 0.2, 11, and 100 mg/m3 or 0.1, 8, and 82 mg/m3, re-

side vignels data for human risk assessment. A simulated con-

sidered occupational exposure monitoring study was conducted to determine typical

breathing zone aerosol concentrations during product use. In the 13-wk study, no
treatment-related changes in daily clinical observations, functional tests, ophthal-
mology, urinalysis, hematology, clinical chemistry, or coagulation parameters were

traces of bromide from the diet to humans, the NOAELs for reproductive and parental effects were 48 and 12 mg

changes. Repeated oral dosing also causes a hypothyroid effect that is specific to rats

been developed elsewhere. Repeated oral exposure in various mammalian species is

inhalation of hexavalent chromium [Cr(VI)] has been associated with increased

exposure and bladder cancer risk, and whether meta relative risks (mRR) differed

bladder cancer. Differences in meta relative risks were compared to RRs predicted by

previous estimate (mRR = 1.11; 95% CI: 0.95–1.30), with no significant association

Arsenic (As) exposure and bladder cancer risk, and whether meta relative risks (mRR)

based on data from southwestern Taiwan, form the basis of the U.S. Environmental

exposure and bladder cancer risk, and whether meta relative risks (mRR) differed

with two recent studies to further examine the association between low-level arsenic

pins as other than drinking water over the past century. To the best of our knowledge,

factors. The resulting RfD range is protective against diffuse hyperplasia,

range of 8 RfD values (2 modeling approaches

metrics together with corresponding incidences for diffuse hyperplasia in each in-

used to predict internal dose measures for chromium in the duodenum, jejunum,

Intestinal tumors have been observed in mice (but not rats) following chronic ex-

ing, specifically villosus blunting and crypt hyperplasia—collectively termed
diffuse hyperplasia. Recent mode of action studies support that these tumors were

Indeed the result of chronic wounding and regenerative hyperplasia to repair the
intestinal mucosa. Herein, we develop an oral reference dose (RfD) that is protective of the tissue parenchyma lesion (diffuse hyperplasia), and therefore is protective of ins-
testinal cancer. A rodent physiologically based pharmacokinetic (PBPK) model was

used to predict internal dose measures for chromium in the duodenum, jejunum,

lumen of mice under the conditions of the 2-year bioassay. These internal dose

metrics together with corresponding incidences for diffuse hyperplasia in each in-
testinal segment were used to characterize the dose-response relationship for the

small intestine in a single plot containing a robust dataset with as many as 24 data

Points of departures (PODs) were derived using benchmark dose modeling and
global nonlinear regression, with models providing acceptable fits differing <3-

Human equivalent lifetime average dose values were estimated for each POD

using two different methods of extrapolation with the human PBPK model for
cromium. Dividing the PODs by uncertainty factors (UFs) of 10–30 yields a
range of 8 RfD values (2 modeling approaches × 2 human equivalent dose methods × 2 UF values). The resulting RfD range is protective against diffuse hyperplasia, and

is therefore protective of both noncancer and cancer effects in the small intes-
tine. This range of RfD values leads to acceptable Cr(VI) concentrations in drink-
ing water that are greater than those typically found in drinking water sources (<5

Inhalation of hexavalent chromium [Cr(VI)] has been associated with increased lung cancer risk among workers of certain industries, but no well recognized or
published mode of action (MOA) exists. Although it has been suggested that
Cr(VI) acts by a mutagenic MOA because it damages DNA in vitro and in some in vivo tests, several recent reviews have concluded that Cr(VI) is only weakly mutagenic and that genetic/epigenetic changes resulting in genomic and/or microsatellite instability, inflammation, oxidative stress and deregulation of repair mechanisms play a role in carcinogenicity. Further, recent data for mouse small intestinal cancers caused by Cr(VI) in drinking water support a cytotoxic MOA involving chronic inflammation and epithelial cell death. Using the dose-response paradigm, we have viewed kinetic, human, animal, and mechanistic data to develop a lung cancer MOA: evaluated plausibility, dose-response, and temporal concordance; and considered alternative MOAs. Among workers with an observed increase in lung cancer, respiratory tissue irritation and inflammation were common clinical findings, and in animal studies irritation and inflammation precede tumor formation in dose (≥50 μg/m3) and time (within 30 days), suggesting that cytotoxicity and inflammation are early key events. The overall evidence supports that Cr(VI) induces lung cancer by a non-mutagenic MOA involving oxidative stress, cytotoxicity, and inflammation, causing oxidative DNA damage, epigenetic DNA modifications and genomic instability, occurring at the high exposure concentrations. Further, extra-cellular Cr(VI) reduction in the lung limits absorption and may introduce non-linearity in the extrapolation of tissue dose from high to low exposures. Based on these findings, the risk assessment for inhalation exposure to Cr(VI) should consider intensity-based dose metrics and non-linear low dose extrapolation approaches.

2237 A Screening Level Assessment of the Health and Environmental Hazards of Organohalogen Flame Retardants.

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Organohalogen flame retardants are extensively used in both industrial and consumer products. However, relatively little is known about the potential of many of these chemicals to cause adverse health and environmental effects. To address this, we conducted a health and environmental hazard screening of almost 100 brominated or chlorinated flame retardants based on the GreenScreen® or Quick Chemical Assessment Tool (QCAT®) methodologies. Priority consideration was given to human health hazards such as carcinogenicity (including mutagenicity and genetic toxicity), reproductive or developmental toxicity, endocrine disruption, and acute mammalian toxicity. Environmental hazards considered included acute aquatic toxicity, persistence, and bioaccumulation. Using publicly available information, each hazard category was assigned a concern level (low, moderate, high, or very high) based on pre-defined numerical ranges, such as no-observed adverse effect levels, and hazard classification schemes from authoritative sources, when available. Less than 10% of the screened chemicals had empirical data to assess each priority hazard category. Where empirical data were not identified, structure activity relationship (SAR) models were relied upon to predict hazard potential. After assigning concern levels for each priority health effect, each chemical received a score, similar to a report card (A, B, C, D, or F). The majority of the screened chemicals received either a D or F grade due to empirical data suggesting high hazard, SAR model predictions, and/or excessive data gaps. Carcinogenicity was the most prominent potential health hazard identified based on empirical data. The most prevalent data gap was endocrine disruption due to the lack of identified empirical data or computer models able to predict this hazard. This study highlights the limited toxicity information available for these widely used chemicals, and indicates that more testing and oversight is critically needed to identify safer alternatives for fire prevention.

2238 Air Quality and Human Health Risks Along the Texas Gulf Coast.

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There are several urban and industrial regions along the Texas Gulf Coast that house some of the largest ports in the US, numerous refineries, thousands of chemical/petrochemical facilities and manufacturing plants. Given its dense population and industrial activity, monitoring and regulation of air quality in these regions is critical for protection of human health. The TCEQ has developed the largest ambient air monitoring network in the US to measure cumulative emissions in the area. Critical areas of concern include areas of high industrial activity, monitoring and regulation of air quality in these regions is critical for protection of human health.
outcomes which differ from those outlined above and are often even more conserva-
tive. This paper explores the background to these divergences and highlights that the
context under which an RfD or DNEL is developed has a significant bearing on the final
determination made.

2241 Risk Estimates for Dioxins, Furans, and PCBs in Edible Fish and Shellfish at the San Jacinto River Waste Pits, Texas, USA.
E. S. Williams, S. Usenko and B. W. Brooks, Environmental Science, Baylor University, Waco, TX.

From the 1960s to the early 1980s, paper process wastes were disposed of at a pitsite on the San Jacinto River. Eventually the pits subsided into the river, spreading polychlorinated dioxins, furans, and biphensyls into the adjacent aquatic ecosystem. Consumption advisories have been in place for local fish and shellfish since 1990. The San Jacinto River Waste Pits site was identified during a Total Maximum Daily Loads process, and was added to the National Priorities List in 2008. Previous ef-
forts to characterize bioaccumulation of chlorinated compounds on the site included sampling of sediment and thirteen species of fish and shellfish. During the sampling period, numerous persons were observed fishing on or near the site. Important fish and shellfish consumed from the area appear to be clams (Mercenaria spp.), blue crabs (Callinectes sapidus), and black drum (Pogonias
cromis). Mean TCDD concentrations in clams, crabs, and fish from the site were 28, 3.6, and 2.25 pg/g wet weight, respectively. TCDF was the only other dioxin/furan congener detected, but ten dioxin-like PCBs were observed in these samples. Excess lifetime cancer risk estimates posed by consumption of contami-
nated fish and shellfish were calculated using default exposure parameters; risk esti-
mates for reasonable maximum exposure ranged from 1.2 × 10−5 to 1.5 × 10−4. Risks in excess of 10−4 under these parameters are associated only with consump-
tion of fish and shellfish consumed frequently and at high exposure frequency. While seasonal fluctuation in consumption patterns is expected, low observed adverse effect level of 407 mg/kg/day established from a subchronic
animal feeding study in rats, and a composite uncertainty factor of 3000 to calculate the RfD of 0.12 mg/kg/day for isooctane. Based on this value, a total allowable concentra-
tion of 840 μg/L was derived. This drinking water action level is protective of pub-
lic health since it used the more toxic n-hexane to fill the data gap, and has a mar-
gin of exposure of over 15,000.

2244 Deriving Dermal Safe Harbor Levels for DEHP Relevant to Consumer Products for Adults.
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Di(2-ethylhexyl)phthalate (DEHP), a plasticizer in various consumer products, is
listed as a chemical known to the State of California to cause cancer and reproduc-
tive toxicity. The Proposition 65 Safe Harbor Levels (SHL) for DEHP include a Maximum Allowable Dose (MADL) for DEHP specific to the oral route, of 410 μg/day for adults (58 μg/day for infants, 20 μg/day for neonatal boys), and no Safety
Handling Level (NSRL) for cancer of 310 μg/day, while not route-specific, was based on an oral study. While DEHP in children's toys is of concern due to mouthing behavior, most 60-day notices of violation for DEHP are related to consumer products intended for adults. Thus, there is a need for realistic, der-
mal-specific SHLs and accurate calculations of exposure. From studies on dermal absorption of DEHP from plastic film (present at 25.5 mg/cm² of film, or 40.37% w/w) applied to the skin of rats (used as a conservative estimate of migration of DEHP from a product into the human body since rats have a higher dermal ab-
sorption than humans), a maximum dermal absorption of 0.1% was determined. Comparatively, the reported oral bioavailability of DEHP is 25%. Using these val-
ues, the dermal MADL would correspond to 102,500 μg/kg/day for adults (410 μg/kg/day x 0.25 oral absorption fraction/0.001 dermal absorption) and the dermal
NSRL would be 77,500 μg/kg/day. From the same study, a dermal absorption rate of DEHP from the plastic was calculated to be approximately 0.24 μg/cm²/hour for the rat which corresponds to 0.016 μg/cm²/hour for humans, given observed
species differences. This can be used to calculate dermal exposures with cor-
rection for differing concentrations. Also, a migration rate of 1.4 μg/cm²/hour can be used to determine the DEHP skin loading for estimating transfer of DEHP from a consumer product onto the hands, and then directly and indirectly to the mouth for comparison to the oral SHLs. In summary, route-specific SHLs are con-
sidered useful for realistically estimating risks associated with exposure of adults to
DEHP from consumer products.

2245 Microfluidic Steroidogenesis Assays for In Vitro Toxicant Screening.
A. B. Theberge 1, 2, 3, E. Berthier 4, C. J. Hedman 5, B. P. Casavant 4, N. P. Keller 1, 4, W. A. Bushman 1 and D. J. Beebe 1, 2. Molecular and Environmental Toxicology, University of Wisconsin-Madison, Madison, WI; 2Biomedical Engineering, University of Wisconsin-Madison, Madison, WI; 3Urology, University of Wisconsin-Madison, Madison, WI; 4Medical Microbiology and Immunology, University of Wisconsin-Madison, Madison, WI; 5Institute for Clinical and Translational Research, University of Wisconsin-Madison, Madison, WI. Sponsor: C. Bradford.

Many endocrine disrupting compounds act by interrupting steroidogenesis, thus disrupting subsequent processes that rely on steroids for signaling such as hormone-mediated pathways crucial for development. High-throughput in vitro assays are re-
quired to identify such compounds efficiently. Liquid chromatography-mass spec-
trometry (LC-MS) is a gold standard technique for steroid characterization and quantification, but traditional methods for steroid extraction with organic solvents, typically used to remove matrix components prior to LC-MS, require many manual

2243 Oral Risk Assessment for Isooctane—A Class-Based Approach Using the Surrogate n-Hexane.
B. Wang 1, P. Undesser 2 and M. Whittaker 1. 1ToxServices LLC, Washington DC; 2Water Quality Association, Lisle, IL.

Isooctane is a volatile liquid commonly found in gasoline. It is principally released into the environment via the manufacture, use and disposal of products associated with the petroleum industry. Although the most likely route of human exposure to
2246 An Improved In Vitro Method for Determining Chemical Effects on Steroidogenesis Using LC/MS/MS to Monitor Multiple Steroid Hormones Combined with Gene Expression.

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Steroid hormones play vital roles in development. Environmental compounds can impact the steroidogenic pathway producing a wide range of effects. Quantifying multiple steroids from a single biological sample, coupled with gene expression of the enzymes that form them, allows for a better evaluation of these effects. The human adrenal H295R cell line is currently used by the Environmental Protection Agency (EPA) to screen for chemical effects on steroidogenesis by evaluating production of Estradiol (E2) and Testosterone. Following EPA guidelines, 22R-(hydroxycholesterol (22RHC) can be added (20-40 μM) to the culture medium to increase basal production of E2. Addition of 22RHC to the H295R culture medium bypasses steroidogenic acute regulatory protein (StAR). To evaluate the induction mechanism of 22RHC, H295R exposures were conducted using both 22RHC and forskolin, a known inducer of steroidogenesis. 22RHC and forskolin were administered to cells at 0.4, 1, 10, 20 and 40 μM and 0.03, 0.2, 0.3, 1, 3 and 10 μM, respectively for 48 hr. RT-PCR was used to quantify the expression of 9 steroidogenic genes and LC/MS/MS was used to measure hormone levels of 12 steroids. Forskolin was found to upregulate mRNA of CYP11B2, HSD3B2, CYP21A2, CYP17A1, CYP19A1 in a dose-dependent manner. StAR mRNA was also increased at the highest exposure concentration. No significant upregulation of mRNA was observed in the 22RHC-exposed cells within the solubility limit. Steroid levels from LC/MS/MS displayed first and zero order enzyme kinetics. LC/MS/MS results showed that 22RHC exposures significantly increased early event markers (e.g. progesterone, pregnenolone, DOC, and 11 DOC), whereas forskolin exposures resulted in higher levels of most steroids. These results demonstrate that steroid genes are not regulated by a positive feedback loop in 22RHC-exposed cells. This study shows that evaluation of multiple steroid levels combined with gene expression provides a more complete picture of steroidogenesis.

2247 Mapping Pathways of Endocrine Disruption in MCF-7 Cells: 2D and 3D Culture Systems.

M. M. Vantangoli, S. Hall and K. Boekelheide. Brown University, Providence, RI.

It is becoming increasingly important to identify estrogenic-active compounds early in toxicity testing, placing weight on the use of in-vitro screening. Currently, the pathways of endocrine disruption are being mapped in response to estrogenic compounds. In this study, human breast adenocarcinoma MCF-7 cells are being utilized to map pathways of estrogenic disruption following exposure to estrogenic estrogenic-active compounds. Our study utilizes two models, a 2-dimensional system representative of classical in-vitro studies, and a scaffold-free 3D culture system. Both systems follow a protocol that includes 72 hours of growth in media containing 10% FBS for 72 hours, followed by 48 hours in media containing 5% stripped serum. Treatment conditions of interest begin following the 48 hours. Preliminary data shows that our 2D system is responsive to estrogenic compounds including estradiol, DES and BPA across several endpoints, including cell proliferation and protein expression. Over a range of concentrations of estradiol (0.01nmM-100mM) and times (1, 3, 6, and 12 hours) analysis of gene expression showed increases in classical estrogen responsive genes including progesterone receptor (1.5-3.5 fold), cathepsin D (1.5-2 fold), estrogen-inducible pS2 (1.5-4 fold) and GREB1 (4-14 fold) at concentrations as low as 0.01nmM estradiol. These results were also shown by Western blotting, with time-dependent increases in pS2 and GPR30 following estradiol stimulation. Our 3D system exhibits cell-contact and matrix-dependent responses that may play a role in evaluating toxicities of estrogenic-active compounds. Here, we show the potential use of both 2D and 3D culture of MCF-7 cells to assess the estrogenic activities of suspected estrogenic-active compounds.

2248 Cell-Specific Control of Estrogen Response Mechanisms in HeLa-9903 and T47D-KBgluc Cell Lines.


Estrogen signaling can be adversely affected by endocrine disrupting chemicals via agonism or antagonism. Cell lines capable of detecting agonism and antagonism are useful screening tools for determining estrogenic activity of chemicals. Transcriptional effects of the Estrogen Receptor (ER) are modulated by interactions with coregulatory proteins that function as either coactivators (agonism) or corepressors (antagonism). Selective ER modulators (SERMs), such as tamoxifen, favor recruitment of coactivators that inhibit transcriptional activity. In different tissues, tamoxifen can have partial-agonist-antagonist activities which may be related in part to the milieu of ER coactivators and corepressors in these tissues. To further elucidate effects of SERMs on transcriptional activity, agonism and antagonism was evaluated using reporter gene assays in two cell types, hERα-HeLa-9903, derived from a human cervical tumor, and T47D-KBgluc, stably transfected breast cancer cells. Cells were acclimated in 96-well plates and exposed for 24 hr to tamoxifen (TAM), 4-OH tamoxifen (4HTAM), raloxifene (RALON), estradiol (E2), dihydrotestosterone (DHT), corticosterone (CORT), and 17β-estradiol (EST). Cytoxicity, luminescence, and solubility were measured. An agonist response was demonstrated for TAM, RALOX and 4HTAM in hERα-HeLa-9903 cells, but not T47D-KBgluc. Antagonism also showed distinct results for the two cell types where T47D-KBgluc demonstrated a potential additive effect with 4-HTAM and the maximal response was 4-fold higher than E2. RALOX and TAM also demonstrated enhanced responses in hERα-HeLa-9903 cells. Because T47D-KBgluc cells have endogenous ERα and ERβ, and hERα-HeLa-9903 cells have exogenous ERα, coupled with the fact that coactivators and corepressors may be different in breast and cervix tissues, the resulting endocrines control agonism and antagonism may be regulated, at least partially, by separate and distinct gene transcriptional pathways. These findings show that measuring estrogenic activity in two cell lines may provide a more accurate assessment of estrogen response.

2249 Profiling of ERα-Coregulator Binding As a Means for Functional Classification of Unknown Endocrine Disruptors.

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Testing chemicals for their endocrine-disrupting potential, e.g. interference with estrogen receptor alpha (ERα) signaling, is an important aspect of chemical safety profiling. Due to drawbacks of in vivo testing, the development of in vitro alternatives has a high priority. In a previous study, we have demonstrated an in vitro assay which profiles binding of (un)liganded nuclear receptors to a microarray of coregulator-derived peptides, as a good candidate. Here, a set of 13 model compounds, US-EPA recommended for ERα gene reporter assay proficiency testing, was used to assess reproducibility, robustness and added value of our assay. ERα-coregulator binding profiles in the presence of a concentration series of each compound were generated. With a median coefficient of variation of 5% and excellent correlation (R2=0.99) between replicate measurements, the unsaturation level of the peptide microarray was well within the range observed for other compounds used in vitro ER functional assays. Per compound, a dose-response curve for each ERα-coregulator interaction was constructed. Our results show correct prediction of estrogenicity for 12 out of 13 tested compounds. The potency ranking for 9 out of the 10 ERα agonists exactly matched that for transcriptional activation as reported by ICCVAM and US-NIEHS with excellent correlation (R2=0.98). Moreover, unsupervised classification (Hierarchical clustering, Euclidean distance, average linkage) with the compound-characteristic ERα-coregulator binding profiles results in structurally related compounds to cluster together, whereas the steroid test compounds having an aromatic A-ring were separated from those with a cyclohexene A-ring. This latter feature should be exploited to build a prediction model to enable classification of unknown toxicants by comparison with pre-profiled references.
Endocrine disruptor compounds (EDCs) are a group of natural or synthetic compounds that have the capacity to interact with the endocrine system of living organisms and consequently causes adverse health effects in an intact organism, or its progeny, or (sub)populations. Due to the impact that this interaction could have on human health, there is an increasing interest in assessing the risk of the exposure to EDCs. Currently, several in vitro and in vivo assays have been developed and few of them validated and regulated accepted. For instance, the US EPA developed the Endocrine Disruptor Screening Program, which has been recently implemented. For the program a large number of experimental animals will be still use even for testing some of the in vitro assays. Herein, we performed the inter-laboratory validation of two robust models that addresses agonistic and antagonist effects of the human hormone receptor, the YES (Yeast Estrogen Screen) and the YAS (Yeast Androgen Screen). Both assays are non-animals alternatives to the estrogen and androgen receptor binding assays proposed in the EDSP and OECD Conceptual Framework. The ring trial is the final experimental part of the validation process at the European Center for Validation of Alternative Methods (ECVAM). A set of 24 blinded compounds (7 estrogens and 6 androgens of diverse potency, 3 anti-estrogens, 3 anti-androge and 5 negative compounds) have been tested in five different laboratories. The analysis of the first phase of this ring-trial already demonstrates a high reproducibility for both methods among the different participating laboratories.

NICEATM conducted an international validation study of the BG1Luc estrogen receptor (ER) transactivation (TA) test method. The test method evaluation report was reviewed and the method accepted by U.S. regulatory agencies and the Organisation for Economic Co-operation and Development. In 2011, NICEATM nominated the BG1Luc ER TA to Tox21 to be evaluated for adaptation into a quantitative high throughput screening (qHTS) assay. The Tox21 collaboration, an effort by the National Toxicology Program, NIH Chemical Genomics Centers, Environmental Protection Agency, and Food and Drug Administration, was formed to advance toxicity testing by shifting from traditional in vivo tests to in vitro methods. A major goal of Tox21 is to prioritize chemicals for in-depth toxicity testing. One method for prioritization is the use of qHTS assays using cell- and biochemical-based assays to construct concentration–response curves for thousands of chemicals. The Tox21 consortium adapted the BG1Luc ER TA manual method to a qHTS format, making it the first assay validated for regulatory use to be adapted to Tox21. Data from qHTS assays have been generated for approximately 10,000 chemicals in both the agonist and antagonist versions of the qHTS assay. Seventy-six chemicals had been tested in both the manual and qHTS methods. Data from both methods were used to evaluate the degree to which classifications of test chemicals in the BG1 manual and qHTS methods matched reference classifications (accuracy) and the degree to which chemical classifications were identical between the two methods (concordance). Except for a few discrepancies attributable to different test concentrations used in the two methods, the BG1Luc ER TA manual and qHTS methods produced almost identical results in terms of accuracy, with a high degree of concordance. Supported by ILS staff under NIEHS contract N01-ES-35504.

Reconstructed Vaginal-Ectocervical and Endocervical Tissue Models for HSV-2, Chlamydia, and Gonorrhea Infections.

Despite extensive efforts, limited success has been achieved in developing tissue models for sexually transmitted infections (STIs) such as herpes simplex virus 2 (HSV-2), Chlamydia trachomatis (Ct), and Neisseria gonorrhoea (Ng). We developed highly differentiated, normal human 3-dimensional (NHu-3D) vaginal-ectocervical (EpiVaginal™) and endocervical (EpiCervical™) tissue models for STI infections. Immunohistochemistry and quantitative real time PCR were used to characterize the ectocervical tissues to monitor HSV-2 infection. Infection of the ectocervical tissue model by the infectious elementary body (EB) of Ct was assessed by quantitative cultures. ELISA assays were used to quantify TNF-α release in response to Ng infection. Results show that EBs infected vaginal tissue express nectin-1, a receptor for HSV-2, and infection experiments showed that the tissue model was infectable with HSV-2. Similar to the in vivo situation, infection of the vaginal-ectocervical tissue model with HSV-2 caused a separation of the epithelium from the lamina propria layer (“blister formation”). The results were confirmed by DNA PCR. The data from Ct infected endocervical tissues showed a cycle level of EBs which were 1) present at 12 hrs, representing bacteria that failed to be taken up by the cells, 2) not detectable at 24 hrs, all intracellular EBs might have been converted to the replicative reticular bodies (RB) and are therefore not cultivable and 3) present at 40 hr, RBs have been converted back to EBs and are now able to induce the first cycle. Furthermore, the reconstructed endocervical tissues respond to Ng infection and TLR ligands by secreting TNF-α into the culture medium. In conclusion, new in vitro reconstructed ectocervical and endocervical tissue models have been developed for HSV-2, Ct, and Ng infections. The models can be used to study the safety and efficacy of candidate therapeutics aimed at preventing or neutralizing HSV-2, Chlamydia, or gonorrheal infections.

Combining Pathway-Based In Vitro Assays to Prioritize 1848 Environmental Chemicals for Estrogenic Potential.

There are thousands of environmental chemicals subject to regulatory decisions for estrogenic potential. Due to the resources required to perform traditional toxicity tests, high-throughput screening (HTS) assays have emerged as a viable tool for chemical prioritization. The ToxCast and Tox21 programs have tested 1848 chemicals in a broad screening panel of 19 assays for estrogen receptor (ER) agonist and antagonist activity. These assays screen for ER activity by profiling effects on ligand binding or cellular changes across a diverse array of assay types (e.g. receptor binding, protein complementation, transcriptional activation, and cell growth) and cell types (T47D, BG1, HEK293T, CHO-K1, HeLa). The protein complementation assays were run with and without S9 to determine metabolic potential. Assays were assigned to assay- groups to distinguish ER mechanisms (i.e. agonism, antagonism, metabolic activation/deactivation, ERa/ERβ activation). Concentration response data for each chemical was normalized to 17βestradiol, scaled for differences in assay sensitivity, and fit to a Hill model. A composite concentration response curve for each chemical-group combination was created from curve fitting parameters to develop a weight-of-evidence metric for estrogenicity. Composite curve efficacy, potency, and “goodness of fit” were used to rank chemicals and distinguish putative partial agonists from discordant assay results. In a separate analysis, uterotrophic assay results (39 chemicals) were used to characterize composite curve potency using logistic regression. Overall, of the original 1848 chemicals 22 were active in at least 13 HTS ER agonist assays, including known ER agonists, 4-nonylphenol, and DES. 347 chemicals were active in at least 3 HTS ER agonist assays, indicating the need for assays measuring multiple biologically plausible ER mechanisms in order to confidently prioritize chemicals for estrogenicity. This abstract does not necessarily reflect Agency policy.

Identification of Compounds That Activate Aryl Hydrocarbon Receptor Using a qHTS Platform.

The basic helix-loop-helix per-Arnt-Sim (bHLH-PAS) superfamily of transcription factors plays an important role in mediating the biological response to endogenous and xenobiotic small molecules. The Aryl hydrocarbon receptor (AhR) is a prototypical member of the bHLH-PAS, and is crucial to responses to environmental changes. AhR mediates cellular responses to environmental pollutants such as aromatic hydrocarbons through induction of phase I and II enzymes but also crosstalks with other nuclear receptor signaling pathways. To identify potential AhR ligands as part of the Tox21 collaboration, we have optimized and miniaturized a cell-based AhR luciferase reporter gene assay in the recombinant human HepG2 cell line Hg27.5c into a 1536-well plate format. We have evaluated this assay by screening a library of 1280 pharmacologically active compounds (LOPAC) plus 88 Tox21 chemicals in triplicate using a quantitative HTS (qHTS) platform.

Identification of Compounds That Activate Aryl Hydrocarbon Receptor Using a qHTS Platform.
From the primary screen, we have identified a group of relatively potent compounds in vitro: CGS-15943, an adenine receptor agonist, 7-HAQ, a H4 receptor antagonist; the cyclin-dependent kinase inhibitor kenpaullone; the amiloride analogue phenamil; and the gastric proton pump inhibitor omeprazole. Validation of the cell-based assay on our integrated robotics system gave a signal to background ratio of 5 and average Z' factors of 0.4, that indicates this assay is suitable for qHTS of the Tox21 10K library. These findings support the utility of a cell-based AhR luciferase assay system for the high-throughput detection of compounds activating the AhR signal transduction pathway. Supported by EPA Interagency Agreement Y3-HG-7026-03.

Without chemicals in a transparent and interpretable manner. HESS and HESS DB are freely provided in the website of NITE (http://www.safe.nite.go.jp/english/kainin/spa/hess-e.html).

Mitochondrial DNA (mtDNA) variations including single nucleotide polymorphisms (SNPs) have been proposed to be involved in idiosyncratic drug reactions. However, current in vitro and in vivo models lack the genetic diversity seen in the human population. Our hypothesis is that different cell strains with distinct mtDNA SNPs may have different mitochondrial bioenergetic profiles and may therefore vary in their response to drug-induced toxicity. Therefore, we used an in vitro system composed of four strains of mouse embryonic fibroblasts (MEFs) with mtDNA polymorphisms. We sequenced mtDNA from embryonic fibroblasts isolated from four mouse strains, C57BL/6J, MOLF/Eij, CZECHII/Eij and PERA/Eij, with the latter two being sequenced for the first time. The bioenergetic profile of the four strains of MEFs was investigated at both passage 3 and 10. Our results showed that there were clear differences among the four strains of MEFs at both passages, with CZECHII/Eij having a lower mitochondrial robustness when compared to C57BL/6J, followed by MOLF/Eij and PERA/Eij. Seven drugs (nelfadonide, ketoconazole, tolcapone, flutamide, tamoxifen, imipramine and troglitazone) known to impair mitochondrial function were tested for their effect on the ATP content of the four strains of MEFs in both glucose- and galactose-containing media. Our results showed that there were strain-dependent differences in the response to some of the drugs. We propose that this model is a useful starting point to study compounds that may cause mitochondrial off-target toxicity in early stages of drug development, thus decreasing the number of experimental animals used.

For the chicken egg genotoxicity assay, an in vitro alternative model for assessing genotoxicity, we used white leghorn chicken (Gallus gallus) egg fertilized before full development of the nervous system. Injections were made on days 9, 10 and 11, the last 3 hours before termination. Livers were harvested for the COMET assay for DNA strand breaks and nucleotide postlabeling for DNA adducts. To deliver test substances, a variety of vehicles were evaluated, covering the range (from hydrophilic to hydrophobic) of solubility, using the endpoint of viability of the embryonic-fetus. The following suitable vehicles were selected, i.e., 50 μl of deionized water, 50 μl of 0.5% aqueous methylcellulose, and 50 μl 20% Solutol HS15 for hydrophilic, amphiphilic, and hydrophobic substances, respectively. Test substances were injected with a 1 ml plastic BD syringe using a 0.4 mm x 13 mm needle. Following the injection into the air sac, the eggs were sealed with paper tape. Test compounds included diethylaminoethanol (hydrophobic) and 2-aminoterephthalamido-cyclohexane (hydrophilic). The eggs were incubated and maintained at 37° ± 0.5°C and 60% ± 5% relative humidity. On day 11, the shells of viable eggs were opened at the blunt end, and the allantochorionic membrane was retracted to allow access to the entire anterior (visceral) aspect of the chicken fetus via the yolk sac. Fetal weights were recorded after removal of the surrounding yolk and tissue. The yolk sac was opened and the entire liver was removed and weighed. Diethylnitrosamine (0.25 - 4.0 mg cumulative dose per egg) and 2-aminoterephthalamido-cyclohexane (0.1-0.6 mg) were tested over a dose range and subsequent alkaline COMET assays were conducted on isolated liver cells. Both compounds gave a positive dose response with plateau occurring at the higher doses. In addition, 2-aminoterephthalamido-cyclohexane was positive in the nucleotide postlabeling assay and showed patterns of DNA adducts similar to those previously observed in turkey eggs and rats.

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in cell index, indicative of cytotoxicity, after exposure to 12.5-50μM DCMBQ over a 72-hour period. Using the alkaline comet assay, a significant genotoxic effect (one-way ANOVA, p<0.05), indicated by increased tail moment, was observed in cells exposed to ≥20μM DCMBQ for 24 hours. To further determine the mechanism of toxicity, reactive oxygen species (ROS) production was measured using the fluorescent DCFDA (2’,7’-dichlorofluorescein diacetate) in cells exposed to DCMBQ (5-50μM) for 2-72 hours. Significant time and concentration dependent increase in ROS were observed (two-way ANOVA, p<0.05) for ≥15μM DCMBQ at 24 hours of exposure but as early as 4 hours of exposure for 50μM DCMBQ. This effect was significantly reduced by the addition of N-acetylcysteine (NAC), a ROS scavenger. Likewise, simultaneous addition of NAC to cells treated with high concentrations of DCMBQ in the alkaline comet assay reduced tail moment values to those comparable with untreated control groups. Based on these results, we conclude that DCMBQ is cytotoxic and genotoxic under these experimental conditions, and these effects are due at least in part to ROS production. As DCMBQ has been detected in finished drinking water samples, additional testing is urgently required to determine potential effects on human health.

2260 Intra- and Interlaboratory Validation Studies on Reactive Oxygen Species Assay for Photosafety Evaluation of Pharmaceuticals.

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A reactive oxygen species (ROS) assay was previously developed for photosafety evaluation of pharmaceuticals. Although outcomes from the previous multicenter validation study were indicative of satisfactory transferability, reproducibility, and predictivity of the ROS assay using Atacrit comet CPS/CPS plus solar simulators, the feasibility of different solar simulators for the ROS assay has never been elucidated. Herein, in 4 participating laboratories, 2 standards and 42 coded chemicals, including 23 phototoxins and 19 non-phototoxics were assessed by the ROS assay using Seric SXX-2500V2 solar simulators with the aim of evaluating the compatibility of different solar simulators for the ROS assay. In the ROS assay on quinine (200 μM), a typical phototoxic drug, the intra- and inter-day precisions (coefficient of variation; CV) were found to be 1.7-9.4% and 2.7-6.9%, respectively. The inter-laboratory CV for quinine averaged 13.2% for singlet oxygen and 7.1% for superoxide. The ROS assay on 42 coded chemicals (200 μM) provided no false negative predictions as compared to the in vitro/in vivo phototoxicity, although several false positives appeared. These results were regarded a convincing demonstration that the ROS assay is compatible with, and can be adapted to, other available suitable solar simulators without losing performance.

2261 Investigating the Role of Mitochondrial Dysfunction in Zoniporide Toxicity.

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Zoniporide, an inhibitor of the Na+/H+ exchanger-1 was developed for the reduction of myocardial ischemic injury in acute coronary syndromes, in the high-risk surgical setting, and secondary for prevention in patients with ischemic diseases. A 28-days intravenous infusion rat study revealed target organ toxicity of the sciatic nerve, spinal cord, and stomach injury (Petersen et al., 2008). Due to insufficient efficacy, zoniporide was discontinued after the phase 2 clinical trial. We have previously reported that zoniporide seems to affect mitochondrial function (Rana et al., 2011) here, we expand on our studies by investigating further mechanisms that could lead to this mitochondrial disturbance. We tested zoniporide in rat liver mitochondria for its inhibition of mitochondrial respiration and conducted mitochondrial swelling experiments to study possible mitochondrial permeability transition pore (MPT) effects of zoniporide. We further tested this compound in H9c2 cells growing in glucose and galactose media and measured ATP depletion and Caspase 3/7 levels. Finally, we tested zoniporide in THLE cells. Next, we wanted to know if we could rescue the detrimental effects of redox cycling on cell health, by incubation with two different antioxidants. One of them was catalase, which effectively dismantle hydrogen peroxide with water as a byproduct, the other was N-acetyl-cysteine (NAC), which minimizes oxidative stress.

Oxidative stress is one of the major mechanisms of drug induced toxicity. One electron reduction of oxidants generates reactive oxygen species (ROS) via redox cycling. In biological systems, the flavo enzymes mediate the transfer of electrons to the quinone by reducing it to the semiquinone. Recently, we evaluated flavin analogues for their ability to redox cycle and tried to establish their association to toxicity. We investigated menadione (a quinone analogue) and tosloxafin (a flavin analogue) for its attribution to its redox cycling activity that leads to oxidative stress which eventually leads to cell death. In our proof of concept study, we investigated the redox cycling capability of menadione and tosloxafin by utilizing a biochemical assay that measures the H2O2 produced through redox cycling of compounds. Next, we tested these compounds in THLE cells using high content imaging to assess the production of ROS using dihydroethidium (DHE). In addition, we measured glutathione and ATP levels in THLE cells. Next, we wanted to know if we could rescue the detrimental effects of redox cycling on cell health, by incubation with two different antioxidants. One of them was catalase, which effectively dismantle hydrogen peroxide with water as a byproduct, the other was N-acetyl-cysteine (NAC), which minimizes oxidative stress. Here, we report that both, menadione and tosloxafin redox cycle and produce hydroxide peroxide (ROS), both in the biochemical assay as well as in the cell based assay. Glutathione levels were also depleted with both of these compounds. Furthermore, the ROS formation and glutathione depletion lead to cell injury that is measured by ATP depletion. Catalase was able to dismantle H2O2 in the biochemical assay. In the cell based assay, ROS, glutathione and ATP effects were not recovered in the presence of NAC. In summary, we established the involvement of redox cycling of quinones and flavin chemotypes to toxicity in THLE cells.

2262 Validation of an HTS-Amenable Assay to Detect Drug-Induced Mitochondrial Toxicity.

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Drug-induced mitochondrial dysfunction has been shown to contribute to organ toxicity and late stage attrition. Therefore, testing for drug-induced mitochondrial dysfunction pre-clinically is vitally important and has the ability to greatly impact the success of a potential drug candidate. Several assays have been developed but are hampered for high-throughput screening because they either require special reagents, isolated mitochondria or specialized equipment.

Here we validate the assay using the classical mitochondrial toxins antimycin A, CCCP and, oligomycin, as well as two drugs known to have mitochondrial liabilities (nenflazine, and flutamide). We tested non-specific detergent digitonin, in a 384-well plate format with a 2 hour exposure. We determined the assay to have excellent reproducibility with less than 3 fold differences between IC50 values form day to day. Once the assay was validated, we screened a set of 75 commercial compounds that included compounds known to cause diffuse organ toxicities and with known or unknown mitochondrial liabilities. Our screening data identified that compounds could be sorted into 7 different categories based on the calculated IC50s of each condition, the cytotoxicity measurement in both glucose and galactose-grown cells and the ATP measurement in both media conditions. Our results are currently evaluated for their sensitivity and specificity in comparison to other existing assays.
A tier I rat primary hepatocyte multi-endpoint cytotoxicity assay (MECA) system is an effective tool to assess and rank order compounds on predicted potential to induce toxicity in repeat-dose in vivo rat toxicology studies. Our MECA system utilizes three primary biochemical endpoint assays: an ATP assay for cell viability, a lactate dehydrogenase (LDH) assay for membrane integrity and a WST-1 assay for mitochondrial function. A positive signal in the WST-1 assay should be further characterized with a Tier II specific assessment of sub-mitochondrial toxicity. To determine potential mitochondrial liabilities, our Tier II mechanistic assessment includes a JC-1 assay for Mitochondrial Pore Transition (MPT) status, a 2’,7’-dichlorofluorescein diacetate (DCFDA) assay for reactive oxygen species (ROS) production, a Caspase 3/7 assay for apoptosis, and a Seahorse XF24™ assay system to determine functional oxidative phosphorylation (mitochondrial respiration), HMG-CoA reductase inhibitors (statins) are compounds that lower cholesterol and reduce cardiovascular disease. There has been increasing evidence that some statins may affect mitochondrial function, leading to adverse effects such as myopathy or rhabdomyolysis. We investigated the effects of multiple statins, including simvastatin, on various mitochondrial functional endpoints. Tier I screening of the statins in primary rat hepatocytes identified a general mitochondrial liability with a 50% decrease in WST-1 throughout the concentration range. In Tier II assays demonstrated that simvastatin treatment induced hepatocellular apoptosis at high concentrations and lowered both basal mitochondrial respiration and maximal respiration compared to control at non-cytotoxic concentrations. These data corroborate the decreased WST-1 reading and suggesting impaired electron transport chain function. A multi-tiered approach is essential to fully characterize potential mitochondrial dysfunction and add predictive value to in vitro toxicology screens in early phases of drug discovery.
Trichloroethylene (TCE) is an industrial solvent and a common drinking water contaminant. Previous studies have identified the TCE metabolite DCVC as responsible for increased kidney toxicity and cancer, yet the molecular events mediating renal toxicity an cancer remain controversial. Our studies in yeast provide a foundation for identifying potential mechanisms of TCE renal toxicity and to establish an alternative model for identifying mechanisms of toxicity in humans. A functional genomics approach in yeast identified DNA damage and repair pathways important in response to DCVC exposure. Specifically, mutagenic translesion synthesis (TLS) and nucleotide excision repair (NER) pathways were found to play important roles in DCVC toxicity. This data suggests DCVC may cause direct DNA damage that elicits a mutagenic repair response. Follow-up studies were conducted in the DT40 avian cell to assess if the mutagenic DNA repair response is conserved in higher eukaryotes. The viability of DNA repair mutants was significantly decreased particularly for TLS and NER mutants. Furthermore, western blot analysis showed initiation of TLS repair after DCVC exposure. These results support a conserved DNA damage and mutagenic repair mechanism mediating DCVC renal toxicity. Additionally, the results support a functional genomics approach in yeast as a viable model for identifying mechanisms of toxicity in higher organisms, including humans.

**2272 The Value of Pharmacological Data to Support Systemic Acute Toxicity Evaluation of Compounds: A Profile Analysis.**


Developing alternatives in the area of acute systemic toxicity implies combination of multiple parameters. We showed that integration of cell-death data, pharmacological profiles and physico-chemical properties resulted in a significant improvement of the LD50 prediction model originally developed by CeTox. In order to reduce the false negative rate, correcting factors were applied to the model; this included considerations of a clear dose-response effect, the magnitude of the response and the number of receptors responding. A decrease of the LD50 parameter was expected the cited criteria were met. At a LD50 threshold of 500 mg/kg, the predictive performances of the so-called V1 model were extremely encouraging with an overall concordance of 88%.

The purpose of the study is to get a better understanding of the data generated on the selection of 15 CNS and Heart receptors potentially playing a causative role in the toxic effect. The analysis of the profiles observed with the set of 73 public domain chemicals was completed as follows:

- Comparison of the LD50 values estimated via the different models (V0, V1, experimental)
- Toxicological categorization based upon the LD50 threshold of 500 mg/kg
- Identification of the receptors most responding and understanding of their biological relevance

No variation of the LD50 values was observed for 43 chemicals while an increase or a decrease was assigned to 10 and 20 chemicals respectively. As expected, the decrease in the LD50 value made the model more predictive for 16 out of the 20 compounds. With regards to the toxic compounds, the receptors most responding were a subset of 5 CNS and heart receptors (M2, alpha2, beta1, GABA, N neuronal). Next step will consist of checking the relevance of such receptors for a set of proprietary chemicals. As such, we could envisage the development of a more economical and pragmatic model that would be useful for early screening purposes.

*SOT 2010, SOT 2011, SOT 2012

**2273 Development of In Vitro Organtotypic 3D Epithelial Models for High-Throughput Screening of Toxicological, Immunological and Developmental Signaling Pathways.**

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Currently, there is a growing need for moderate to high-throughput toxicological assays that provide mechanistic data by targeting specific pathways. Here, we describe the development of in vitro organotypic 3D skin (EpiDerm™) and airway (EpiAirway™) models with the added feature of luciferase based transcription factor (TF) reporter functions. To produce the models, early passage normal human epidermal keratinocytes and tracheal epithelial cells were transduced with lentiviral vectors containing the TF response elements linked to luciferase. Stably transduced cells were selected by puromycin resistance, expanded several passages and cryopreserved to produce large pools of cells for organotypic model production. To date, reporters for 6 stress response pathways, including oxidative stress, DNA damage, metal stress, MAPK/inflammation, NFκB and xenobiotic stress have been developed in both the EpiDerm™ and EpiAirway™ models. Each model has demonstrated a dose response to positive control test articles including, TBHQ, Nbutin-3, ZnCl2, PMA, TNFα and TCDD, respectively, with an average induction of 3-15 fold over vehicle when luciferase activity in tissue...
extracts was quantified using a microplate luminometer. TF reporters can be assem-"}

"blished in custom 96-well arrays for screening of unknown test compounds and mon-"}

"itoring of cell signaling pathway activity. Using this format, 3 test articles and a negi-"}

"ve control (N=3) can be tested against 8 stress pathways in a single assay resulting in"}

"a heat map-like profile of pathway activation. The ultimate goal of the project is to"}

"develop a panel of 14 TF reporter models for both skin and airway models that can be"}

"assessed in order to create 96-well high-throughput arrays as well as additional in-"}

"dividual tissue formats. Results from these initial 6 models indicate that EpiDerm™ and"}

"EpiAirway™ reporter models will provide novel tools for conducting mechanistic hu-"}

"man toxicological studies.

**2274 In Vitro Predictive Toxicology for Breast Cancer.**

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Identifying chemicals that increase breast cancer (BC) risk could help prevent BC, but rodent bioassays are expensive. In vitro and computational methods are being developed to predict adverse effects with limited in vivo testing. The present work aims to develop in vitro methods to predict chemicals that can increase BC risk. Animal and human studies suggest that both genotoxic carcinogens and certain hormone exposures increase BC risk. Our initial goal is to identify in vitro tests that predict mammary gland carcinogens (MCs) in rodent bioassays, since few breast carcinogens have been studied in humans. We have previously identified 208 chemicals causing mammary tumors in rodents in at least one study. We used the Chemical Carcinogenesis Research Information System (CCRIS) to compare genotoxicity profiles for MCs with 27 ‘non-carcinogens’ (nonCs) that did not show increased tumors at any site in National Toxicology Program bioassays. We included five assay types: bacterial mutagenicity, and in vitro and in vivo micronuclei and chromosomal aberration. Data from at least one of these were available for 158 MCs and 22 nonCs. We found that most MCs are genotoxic: 87% were positive in >15% of CCRIS entries within any assay type, with or without metabolic activation (MA), while 10% were consistently negative (i.e., no study was positive). In comparison, 59% of nonCs were consistently negative. Since many in vitro predictive toxicity programs do not include MA, we also evaluated whether MA was needed for genotoxicity. Without MA, the MCs that were consistently negative for genotoxicity increased from 10% to 24%, and the number consistently positive (i.e., positive in >15% of entries in every assay type) dropped from 60% to 48%. In conclusion, a high percentage of MCs are consistently genotoxic but 10% are negative in all tests. Without activation almost 25% of the MCs would be consistently negative. The relationship between genotoxicity and cancer has been extensively investigated in all tests. Without activation almost 25% of the MCs would be consistently negative, but not in vivo. PPARα was activated by PFOA and acetachlor. Triclosan was positive in vivo, but not in vitro. None of the chemicals altered serum MCP-1 in vivo (2,5-PCDA and simazine, were positive in vitro). Carbaylb and bisphenol A induced H2AX phosphorylation in vitro but were not positive in vivo. Bisphenol A, acetylcholine and bisphenol A showed anti-androgenic activity in vitro and in vivo. While negative in vitro, 2,5-PCDA, PFOA, carbaylb and triclosan altered androgen re-"}

"sponse profiles to androgen in vivo. Aromatic, aquatic, and oxidative stress in vitro but not in vivo. In contrast, triclosan induced oxidative stress responsive genes in vivo but not in vitro. PPARα pathway activation has a good concordance in vitro and in vivo. Results for anti-androgenicity, MCP-1 and oxidative stress were much less concordant between in vitro and in vivo. This abstract does not necessarily reflect the policies or views of NIH or the USEPA.

**2276 The In Vitro ST3 Neutral Red Uptake Phototoxicity Test: What to Do with UVB Absorbers?**

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The literature suggests and conventional wisdom holds that Ultraviolet B (UVB) is highly cytotoxic to the Balb C 3T3 fibroblast cell line used in the OECD 432 3T3 NRU Phototoxicity Test and thus the assay is not appropriate for test materials that absorb primarily in the UVB portion of the spectrum. The choice of an appropriate light source and filtering is always a critical factor in this or any assay, and, while this Guidance recommends an Ultraviolet A (UVA) dose of 5 J/cm², the UVB dose (if any) or limit of exposure is not addressed. To define the sensitivity of the cells to UVB and evaluate if indeed their sensitivity precluded the use of UVB, we evaluated the Guidance-required cell viability, OD400 absorption, and other endpoints with increased UVB (290 – 300 nm) exposure by removing the tissue culture plate lid during exposure using a xenon arc solar simulator. This direct (uncovered) exposure increased the UVB irradiance from approximately 19 mJ/cm² to approximately 32 mJ/cm², concomitant with the recommended 5 J/cm² UVA dose. While cell survival was modestly reduced, the resulting IC50, OD400 absorption, PIF, and MPE endpoints from this enhanced UVB exposure alone and response to Chlorpromazine indicate no adverse effect on the validity of the assay and accept-"}

"able Guidance-defined results under these exposure conditions. This additional ability to test UVB absorbers provides all the advantages of the assay to this subset of test materials, enhances this step in the preclinical process of drug discovery and further establishes the robustness of the assay as a valid step in preclinical drug de-"}

"velopment.

**2277 Comparison of Multiple Assay Formats to Measure Kinase Inhibitor Activity.**

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Protein phosphorylation is a key mechanism for controlling cellular functions, and protein kinase inhibitors show good efficacy as oncology therapeutics. Currently there are 15 marketed kinase inhibitors, with many more in clinical trials. A key component of drug discovery efforts for these inhibitors is the optimization of kinase selectivity, since a greater degree of promiscuity is associated with a greater risk of safety concerns. A wide variety of biochemical and cellular kinase assays which utilize multiple technologies are available for testing compounds and determining kinase activity, and an important consideration is the translation across these assays. A set of 14 Pfizer kinase inhibitors with different primary kinase targets were pro-"}

"filed across three different types of assays: biochemical kinase assays measuring peptide substrate phosphorylation; ActivX technology, which measures specific binding to kinases within a cell lysate; and cell-based kinase assays, measuring endogenous substrate phosphorylation. The biochemical assays were conducted at Km levels of ATP with nonphysiological peptide substrates, while the cellular assays utilized intact cells and cellular levels of ATP. A comparison of results across these platforms against three different kinases, aurora, abl or glycogen synthase kinase beta, was carried out and the results demonstrate a high degree of congruency with the cellular kinase activity and the ActivX technology. For this set of compounds, there was less agreement between the biochemical versus cellular kinase assays. These results suggest that utilization of high throughput biochemical assays can be used to provide an initial determination of kinase activity, but that followup testing under more physiological conditions using the ActivX technology, or cellular assays, is required to fully assess kinase inhibitor activity.

**2278 Mouse Cecal Microbiota Converts Monomethy larsonic Acid (MMA) to an Array of Ox-"}

"y- and Thio-Arsenical Metabolites.**


The metabolism of arsenicals markedly affects their tissue distribution and reten-"}

"tion as well as the toxic and carcinogenic effects of this metalloid. Metabolism of ar-"}

"senicals by the microbiota of the gastrointestinal tract has been shown to convert inorganic and dimethylated arsenicals to various methylated species. Here, anaero-"}

"bic microbiota from ceca of adult female C57BL/6 mice was incubated with 20,
200, 1000, or 2000 parts per billion (ppb) MMA for up to 48 hours at 37°C. Samples of supernates from reaction mixtures were taken for arsenic speciation by HPLC-ICP-MS. MMA was converted to monomethylated mono- (MMMTA), di- (DMDTA), and tri- (MMTTA) thiolated species. In addition, MMA was also converted to dimethylated mono- (DMMTA) and di- (DMDTA) thiolated species. After 48 hours, DMDTA was the predominant metabolite in reaction mixtures containing 20, 200, or 1000 ppb MMA. In reaction mixtures containing 2000 ppb MMA, MMDTA was the predominant metabolite of MMA. These results show that anaerobic microbiota from mouse cecum includes organisms that efficiently methylate MMA and can convert oxyarsenicals into homologous thioarsenicals. The presumptive source of sulfur used for conversion of oxyarsenicals into thioarsenicals is hydrogen sulfide produced by the microbiota through dissimilatory sulfate reduction. Conversion of methylated oxyarsenicals into methylated thioarsenicals during preabsorptive metabolism may influence the transport of ingested arsenic across the gastrointestinal barrier thereby affecting the systemic distribution, fate, and effects of arsenic. (This abstract does not reflect U.S. EPA policy).

Paraoxonase Activity in Subchronic Low-Level Inorganic Arsenic Exposure.

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Epidemiological evidences indicate close association between inorganic arsenic exposure via drinking water and cardiovascular diseases. While the exact mechanism of this arsenic-mediated increase in cardiovascular risk factors remains enigmatic, studies indicate an oxidative stress role in cardiovascular diseases. To study the association between inorganic arsenic exposure and cardiovascular diseases, rats were exposed to sodium arsenite (50, 100 and 150 ppm) and sodium arsenate (100, 150 and 200 ppm) in their drinking water for 12 weeks. PON activities towards paraoxon (PONase) and phenylacetate (AREase) in plasma, lipoproteins, liver and brain microsomal fractions were determined. Inhibition of PONase and AREase in plasma and HDL characterised the effects of the two arsenicals. While arsenite inhibited PONase by 33% (plasma) and 46% (HDL) respectively, arsenate inhibited the enzyme by 41 and 34% respectively. AREase activity was inhibited by 52 and 48% by arsenite; the inhibition amounted to 72 and 67% respectively by arsenate. The pattern of inhibition in plasma and HDL indicates that arsenite induced a dose-dependent inhibition of PONase whereas arsenate induced a dose-dependent inhibition of AREase. In the VLDL, arsenate inhibited PONase and AREase while arsenite inhibited PONase. In the hepatic and brain microsomal fractions, only the PONase enzyme was inhibited by the two arsenicals. The inhibition was more pronounced in the hepatic microsomes where a 70% inhibition was observed at the highest dose of arsenate. Microsomal cholesterol was increased by the two arsenicals resulting in increased cholesterol/phospholipid ratios. Our findings indicate that decreased PON activity observed in arsenic exposure may be an incipient biochemical event in the cardiovascular effects of arsenic. Modulation of PON activity by arsenic may also be mediated through changes in membrane fluidity brought about by changes in the concentration of cholesterol in the microsomes.

Comparative Early Oncogenic Effects of Cadmium and Arsenic in Human Lung Epithelial Cells.

R. J. Person, E. I. Tokar and M. P. Waalkes. NTP, NIEHS, NIH, Research Triangle Park, NC.

Cadmium (Cd) and inorganic arsenic (iAs) are known human lung carcinogens. In this study, we compare development of in vitro cellular models of human lung cancer induced by Cd or iAs. We have shown chronic, low-level (5 μM) Cd (as CdCl2) induces an acquired cancer phenotype in human lung epithelial cells (HPL-1D) after 20 weeks of continuous exposure. Here we compare an iAs model in development to this previously developed Cd model. The HPL-1D cells that were used are an immortalized, non-tumorigenic human peripheral epithelial cell and were used with both agents. HPL-1D cells were chronically exposed to non-toxic levels of Cd (5 μM) or iAs (2 μM) and over 20–26 weeks of chronic exposure signs of oncogenic transformation were induced. Matrix metalloproteinase (MMP) secretion, colony formation in soft agar, invasion and expression of cancer relevant genes were used to assess oncogenic phenotype in these cell models. By 26 weeks of continuous iAs exposure, secreted MMP-2 significantly increased to 147% of control, near levels typical of a cancer phenotype. In comparison, after only 20 weeks of Cd exposure, MMP-2 levels increased to 358% of control and cell invasion and colony formation increased by more than 5-fold compared to control cells, all indicating an oncogenic phenotype. Following iAs exposure (26 weeks) increases were seen in MT-1A (768% of control) and MT-2A (614% of control) expression, similar to that seen with Cd transformation. Expression of epithelial-to-mesenchymal transition markers (EMT) increased in Cd control with chronic iAs exposure. High VEGF oncogene expression is often seen in lung adenocarcinomas and after 26 weeks of chronic iAs treatment expression increased to 4.3-fold above control. The SLC39A8 (ZIP 8) transporter, which is known to influx Cd, though increased by Cd transformation was unchanged in chronic iAs exposure. Thus, it appears that both Cd and iAs can induce early signs of transformation in human lung epithelial cells, although some gene expression is iAs specific.

Metallothionein Blocks Arsenic-Induced Oxidative DNA Damage.

W. Qu and M. P. Waalkes. NTP, NIEHS, NIH, Research Triangle Park, NC.

Metallothionein (MT) plays an important role in detoxication of inorganics. Inorganic arsenic is a toxic metalloid and a human carcinogen that may act, in part, by causing oxidative DNA damage (ODD). MT can limit ODD induced by other inorganic carcinogens, like cadmium. Although MT can mitigate arsenic toxicity in vitro, how MT impacts arsenic-induced ODD has not been defined. Here, we studied ODD induced by acute arsenic treatment in vitro and the effects of cellular MT using cells that poorly express MT (MT-III double knockout; called MT-null cells) compared to parental wild-type (WT) MT competent cells. MT-null and WT cell lines were first exposed to arsenite (NaAsO2) for 24 h to assess cytotoxicity. Arsenic was much less cytotoxic in WT cells (LC50 = 11.0 ± 1.3 μM, mean ± SEM) than MT-null cells (LC50 = 5.6 ± 1.2 μM). Arsenic-induced ODD was measured by the immuno-spin trapping method which measures DNA radicals after conversion to stable DNA nitrones in situ. Arsenic treatment (1 or 5 μM; 24 h) induced much less ODD in WT cells (121% and 141% of control, respectively) than in MT-null cells (220% and 260%). In WT cells arsenic caused concentration-related increases in MT expression (in transfected and parent) and in metal-activated transcription factor 1 (MTF1), a requirement for induced MT gene expression by arsenic. In contrast, in MT-null cells, the basal levels of MT were very low and were not increased by arsenic. Transfection of MT-I into MT-null cells markedly reduced arsenic-induced ODD. Two important transport genes, Mrp1 and Mrp2, showed increased expression in WT cells but not MT-null cells. Arsenic caused concentration-related increases in the oxidant defense genes, HO-1 and GstA2 in both WT and MT-null cells, but to much higher levels in WT cells. Thus, MT protects against arsenic-induced ODD in MT competent cells potentially by multiple mechanisms including direct sequestration and scavenging oxidant radicals. MT-competent cells are more adept at activating metal transport systems and oxidant response genes, although the role of MT in these responses is unclear.

Increase Blood Pressure, Changes of Left Ventricular Geometry, and Function in Children Environmentally Exposed to Inorganic Arsenic.


Hypertension is a known cardiovascular risk factor to develop final cardiac events and epidemiologic studies in adults have been related hypertension with inorganic arsenic (iAs) exposure. Left ventricular mass (LVM) increase is a potent predictor of cardiovascular morbidity and mortality and it is stimulated by higher blood pressure as well as impaired myocardial contractile performance. Ejection fraction (EF) has been employed as a good index of global systolic function. The aim of this study was evaluate the association between iAs exposure, blood pressure, and echocardiographic parameters in children.In this cross-sectional study 170 children (3-14 years old) chronically exposed to iAs through drinking water were recruited in iAs-endemic area of central part of Mexico. LVM and EF were derived from echocardiographic and blood pressure was measured by standard protocols. Total arsenic in urine (UtAs) was significant associated with systolic (β=2.65; p=0.056) and diastolic blood pressure (β=0.012; p=0.019) blood pressure in multivariate regression models adjusted by age, gender and body mass index. Indeed, diastolic blood pressure was associated with cumulative arsenic exposure (ZAsE; ppb-year) thought drinking water (β=0.003; p=0.002). Notably, 1VM (g) was significant associated with ZAsE (β=-15.43; p=0.002) in adjusted multivariate model. Diastolic prehypertension was present in 48%, concentric remodeling was presented in 60% [normal LVM and relative wall thickness (RWT) ≥0.40] and 43% and LVM concentric hypertrophy defined as LVM >88.9 g/m2 and RWT≥0.43 were presented in 7% (11 children). Moreover, EF was inversely associated with UtAs 35ng/ml (β=-3.21; p=0.036) in adjusted multivariate analyses. In conclusion, iAs exposure would be able to cause diastolic prehypertension, increase of LVM and decrease of EF in children, given evidence of possible cardiovascular disease in early-life exposure.
Dimethylarsinic acid (DMAV) is the major urinary metabolite of inorganic arsenic in humans and most animals. DMAA is weakly cytotoxic, however, it is reduced to dimethylarsinous acid (DMAIII) which is over 100 times more toxic. Although glutathione S-transferase omega 1 (GST01) catalyzes the reduction of DMAAs, its role in DMAA reduction in vivo or in cell extracts is uncertain. We studied the regulation of DMAA reduction in rats and rat liver cytosol to better understand its mechanism. To assess DMAA reduction in rats, we devised a novel procedure. This is based on following the time course of the accumulation in the blood of the RBC-bound dimethylarsenic (DMAa), which represents DMAIII. Therefore, we serially measured the RBC-bound DMAs in the blood of DMAA-injected anesthetized rats with ligated renal pedicles. These studies indicated that reduction of DMAA to DMAIII was rapid, as in 90 min 31% of the injected 50 μmol/kg DMAA dose was converted to DMAIII that was sequestered by the circulating erythrocytes. Pretreatment of rats with glutathione (GSH) depletors (phoron or BSO) delayed the elimination of DMAA and the accumulation of RBC-bound DMAs, whereas the methyltransferase inhibitor PAD was without effect. Reduction of DMAA by rat liver cytosol was assayed by extraction of DMAIII from the incubations of cytosol with DMAA and quantification by HPLC-HG-AFS. We found that reduction of DMAA required cystolic protein and GSH and was inhibited by thiol reagents, GSSG and dehydroascorbate. Although thioridoxin reductase (TrxR) inhibitors (urothiol-glucov and trivalent antimony) inhibited cystolic DMAS reduction, TrxR plus NADPH alone or when added to the cytosol failed to support DMAA reduction. On ultrafiltration of the cytosol through a 3 kDa filter, the reducing activity in the retentate was lost but was largely restored by NADPH. Such experiments also indicated that the reducing enzyme was larger than 100 kDa, and was not GST01. In summary, reduction of DMAA to the supertoxic DMAIII in rats and rat liver cytosol is rapid and GSH dependent, yet its mechanism is still elusive.

**2286** Effects of Treatment with Dimethylarsinous Acid (DMAIII) on the Urinary Bladder Epithelium of Female Arsenic Methyltransferase (As3mt) Knockout Mice and C57Bl/6 Mice.

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Chronic exposure to inorganic arsenic (InAs) is carcinogenic to the human urinary bladder. It produces urothelial cytotoxicity and proliferation in rats and mice. DMAV, a major methylated urinary metabolite of InAs, is a rat bladder carcinogen. DMAIII was shown to be the likely urinary metabolite of DMAV-inducing urothelial changes and is postulated to be one of the active metabolites of InAs. To evaluate potential DMAIII-induced urothelial effects, it was administered to As3mt knockout mice which cannot methylate arsenicals. Female C57Bl/6 wild type and As3mt knockout (group) were administered DMAIII (1.5 μmol/kg) once daily for four weeks. Urothelial effects were evaluated by light and scanning electron microscopy (SEM) and immunohistochemical detection of bromodeoxyuridine (BrdU). DMAIII significantly increased the BrdU labeling index in the knockout group compared to control and to the treated wild type group. DMAIII induced a greater increase in the incidence of simple hyperplasia in knockout mice (4/10) compared to wild type mice (2/10). All treated knockout mice had more and larger intracytoplasmic granules, compared to the treated wild type mice. Changes in SEM classification were not significant. In conclusion, DMAIII induces urothelial toxicity and regenerative hyperplasia in mice and most likely plays a role in inorganic arsenic-induced urothelial changes. However, in mice, DMAV does not induce hyperplasia, suggesting that urinary concentrations of DMAIII do not reach cytotoxic levels in DMAV-treated mice.
Epidemiological studies have shown a strong link between chronic arsenic exposure and bladder cancer. An immortalized human urothelial cell line, UROtsa has been widely used as a model of arsenic-induced bladder toxicity. Chronic exposure to 1μM sodium arsenite transforms UROtsa to a cancerous cell line, URO-ASSC. This phenotype is stable with no further arsenite selection, and URO-ASSC is typically assessed for malignancy without arsenite exposure. In the absence of arsenite, we found that both UROtsa and URO-ASSC demonstrate constitutive autophagy. Recent evidence suggests that autophagy is a survival mechanism for cancer. This led us to hypothesize that disrupting autophagy could reduce malignant potential. We found that both UROtsa and URO-ASSC demonstrate autophagy. Impairing autophagy in URO-ASSC by siRNA against ATG7 resulted in 65% reduction of anchorage-independent growth, a key phenotype of arsenic-induced malignant transformation. Surprisingly, when we reintroduced URO-ASSC cells to 1μM of arsenite exposure, the cells’ growth and morphology were not affected, suggesting a higher rate of autophagic turnover in URO-ASSC. Our findings suggest that disruption of autophagy could reduce malignant potential, a finding with translational impact.

Arsenic is an environmental pollutant that induces apoptosis in various tissues. However, underlying molecular mechanisms of arsenic-induced apoptosis are not clear. Recently, we have shown in vitro and in vivo that arsenic inhibits transcriptional activation of the liver X nuclear receptors (LXR), key regulators of macrophage lipid homeostasis. Here, we evaluated the role of LXRx in arsenic-induced apoptosis. Using the ApoE-/- mouse model, we saw significant changes in plaque composition, and thus in LXRx-dependent manner. In the ApoE-/- mice, we assessed plaque staining of: 1) lipid deposition and macrophage content and 2) collagen composition and smooth muscle cell content. Interestingly, arsenic decreases macrophages in LXRα/ApoE-/- mice, where no change was observed in ApoE-/- exposed mice. However, arsenic increased lipids in both genotypes, suggesting impaired macrophage cholesterol efflux capacity and subsequent lipid accumulation. Second, we observed that arsenic decreased collagen content in LXRα/ApoE-/- and ApoE-/- to the same extent, but arsenic increased smooth muscle cells, a major collagen producing cell type, in the in LXRα/ApoE-/-, while they were decreased in ApoE-/-.

As3MT may be an important determinant of risk associated with chronic iAs exposure. Concentrations of iAs, MAs, and DMAs in urines from residents of Churchill County, Nevada, were used to calculate primary (MAs/iAs) and secondary (DMAs/MAs) methylation indices (MI). As3MT genotypes were determined for 198 individuals selected on the basis of lowest and highest secondary MIs. The incidence of an As3MT polymorphism (rs1191493) that replaces a methionyl residue in position 287 with a threonyl residue (M287T) affected the secondary MI. For the M287T polymorphism, median values for secondary MIs were 6.5 in 150 individuals homozygous for wild-type As3MT, 2.8 in 43 individuals heterozygous for wild-type and mutant alleles, and 2.4 in 5 individuals homozygous for the M287T polymorphism. In contrast, median values for primary MIs were unaffected by this polymorphism. Two intronic variants (T35587C and G53991A) reported to alter the levels of MAs and DMAs in other studies did not alter primary and secondary MIs in this population. These results indicate that the common M287T polymorphism of As3MT is associated with altered profiles of methylated arsenicals in urine from individuals chronically ingesting iAs in drinking water. Linkages among As3MT genotype-dependent alterations in urinary arsenical profiles, the catalytic properties of As3MT variants, and disease susceptibility require further examination. (This abstract does not reflect U.S. EPA policy).
Treatment of HepG2 cells with MMA(V), DMA(V), or TMA(V) alone significantly induced CYP1A1 mRNA, protein, and catalytic activity levels. Furthermore, when the cells were co-exposed to MMA(V), DMA(V), or TMA(V) in the presence of TCDD, there was further potentiation of the TCDD-mediated induction of CYP1A1 mRNA, protein, and catalytic activity levels. In addition, MMA(V), DMA(V), and TMA(V) in the absence and presence of TCDD induced the AhR-dependent XRE-driven luciferase reporter activity, suggesting an AhR-dependent mechanism. In conclusion, this is the first demonstration that As(III) metabolites, MMA(V), DMA(V), and TMA(V) induce CYP1A1 mRNA, protein, and catalytic activity levels in an AhR-dependent mechanism and represents a novel mechanism by which As(III) causes carcinogenicity. Supported by NSERC Discovery Grant RGPIN 250139-12.


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Several methods are used for quantifying the toxic inorganic arsenic (iAs) metabolites, methylarsonic acid (MA(V)) and dimethylarsinic acid (DMA(V)), including reversed-phase high-performance liquid chromatography-inductively coupled plasma-mass spectrometry (HPLC-ICP-MS) and hydride generation-cryotrapping-atomic absorption spectrometry (HG-CT-AAS). While HG-CT-AAS has consistently detected these arsenicals in biological samples, HPLC-ICP-MS has provided contradictory results. Here, we compare the capacities of both methods to detect and quantify MA(V) and DMA(V) in an in vitro methylation system containing recombinant human arsenic (+3 oxidation state) methyltransferase (AS3MT). S-adenosyl methionine, a non-thiol reductant tris(2-carboxyethyl)phosphine, and arsenic trioxide (As2O3) or As(V) as substrates. HPLC separation of the in vitro methylation mixture resulted in significant losses of MA(V) and DMA(V) with total arsenic recoveries below 25%. Ultrafiltration showed that both MA(V) and DMA(V) are bound to AS3MT. Oxidation of the mixture with H2O2, prior to HPLC separation increased arsenic recoveries to 95% but oxidized MA(V) and DMA(V), thus preventing quantification of these metabolites. In contrast, direct HG-CT-AAS analysis revealed large quantities of MA(V) and DMA(V) and high total arsenic recoveries (>72%) after cytostatic treatment. These data suggest that HPLC-ICP-MS can provide false-negative results when used for analysis of MA(V) or DMA(V) in biological samples containing protein at concentrations as low as those commonly found in human urine.

2294 The Retention of Trivalent Arsenic Metabolites in Urothelial Cells Is Associated with Markers of As Exposure and Diabetes.

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Exposure to arsenic (As) is associated with an increased risk of many serious illnesses including type 2 diabetes, cardiovascular disease, and reproductive and developmental problems. Previous research has shown that As can act as an endocrine disruptor, altering the regulation of gene expression by numerous nuclear hormone receptors, including the Glucocorticoid Receptor (GR). At very low doses (0.05-1 μM) As enhanced hormone-mediated, GR-regulated gene expression by 2- to 3-fold. Considering the non-cytotoxic nature of these concentrations (1-5 μM) As inhibits receptor-mediated gene expression. We hypothesized that these differential effects reflect separate mechanisms with different targets and that the inhibited activation could be caused by 1) altered rate of hormone receptor translocation to the nucleus, 2) altered number of receptors that translocate, 3) altered steady-state nuclear levels of receptor (e.g., by decreases in nuclear export), or 4) altered efficiency of receptor function, or some combination of these. To test this, we used HEK293 cells to generate a cell line stably expressing rat GR fused to photo-activatable Green Fluorescent Protein (HEK293-pGFP-rGR), allowing
for intracellular tracking of GR using microscopy. The pAGFP-rGR leaves biochemically in a similar manner as native GR. PaGFP-rGR mediates hormone-induced gene expression and microscopy showed that the rate and extent of nuclear translocation of paGFP-rGR increase in a concentration dependent manner in response to synthetic glucocorticoid. We found that intermediate As concentrations reduce this translocation. (Funded by NIH-NIEHS SRP (P42 ES007373), NSF REU (0115378), and NIH-NRSA (2T3 ES 2727-2)).

2297 In Utero Arsenic Exposure and Epigenetic Changes in the Mouse Liver: Comparisons with Transcriptional Modulation.

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Chronic exposure to high levels of iAs in drinking water is associated with cancers of the skin, urinary bladder, lung, liver, and prostate in humans. Exposure to iAs during fetal development has been of particular concern as this developmental time point is often particularly sensitive to the effects of environmental toxicants. In C3H and CD1 mice, iAs acts as a complete transplacental carcinogen in which male offspring born to pregnant C3H and CD1 females exposed to 85 ppm iAs in drinking water during the latter part of gestation (gestational day (GD) 8-18) have an increased incidence of hepatocellular carcinomas (HCCs) in adulthood, a major form of cancer associated with chronic iAs exposure in humans. Analyses of normal-appearing liver tissue from transplacental-exposed newborn livers and tumors with liver tumors have previously reported perturbations in gene expression and/or global DNA methylation levels and suggested these alterations may contribute to iAs-associated liver carcinogenesis. Here, we examined the gene expression profiles of >35,000 transcripts and the DNA methylation levels of >15,000 CpG islands associated with gene promoters of fetal male CD1 mice (GD 18) transplacently exposed to a hepatocarcinogenic dose (85 ppm) of iAs (GD 8-18). Compared to vehicle-exposed mice, we observed statistically significant changes in the transcriptome (308 transcripts) and DNA methylation (191 gene promoters) in gestationally-exposed fetal males (p<0.05, 1.3-fold change vs. controls). Surprisingly, when using same criteria for the identification of changes in DNA methylation and transcript levels, there were no common genes. These results suggest alterations in the DNA methylation may not necessarily be predictive of changes in gene expression and the relevance of these alterations in disease development warrant further study.

2298 Arsenic Compromises Airway Epithelial Barrier Properties in Primary Mouse and Immortalized Human Cell Cultures.

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Arsenic is a lung toxicant that can lead to respiratory illness through inhalation and ingestion. Lung effects from arsenic exposure include lung cancer and obstructive lung disease, as well as reductions in lung function and immune response. As a key player in innate immune defense, airway epithelial cells provide a barrier that protects underlying tissue from inhaled particulates, pathogens, and toxicants. In animal and human studies, arsenic ingestion can lead to altered lung function suggestive of epithelial barrier dysfunction. In this report, we evaluated the effects of a five-day exposure to environmentally relevant levels of arsenic (i.e., < 4 µM as Na-arsenate; equivalent to ~300 ppb) on airway epithelial barrier properties. In a primary mouse tracheal epithelial (MTE) cell model we found that both micromolar (3.9 µM) and sub-micromolar (0.8 µM) arsenic concentrations reduced transepithelial resistance, a measure of barrier function. Immunofluorescent staining of arsenic-treated MTE cells showed altered localization patterns of barrier proteins claudin-1 and occludin at cell-cell contacts. In order to better quantify arsenic-induced changes in barrier molecular components we used the same arsenic exposure on an immortalized human bronchial epithelial cell line (16HBE14o-). We found that arsenic increased the protein expression of claudins -4, -5, and -7 as well as the mRNA levels of claudin-7 in 16HBE14o- cells. Additionally, micromolar levels of arsenic resulted in altered phosphorylation of occludin. In summary, exposure to environmentally relevant levels of arsenic can alter both the structure and function of airway epithelial barrier constituents and, consequently, basic innate immune defense in the airway.


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To understand associations of single nucleotides polymorphisms (SNPs) and haplotypes in arsenic [3+ oxidation state] methyltransferase (AS3MT) with arsenic metabolism, we investigated local residents from arsenic-contaminated areas of Vietnam. Analysis of 18 SNPs revealed that there were four haplotype (HT) groups (HT1: AS3MT 03963 – 06144 – 12390 – 14215 – 35587 – 37950; HT2: AS3MT 04602 – 35991, HT3; AS3MT 05913 – 09749 – 27215, HT4: AS3MT 358903 – 37853) in this population. Urinary monomethylarsonic acid (MMA)/inorganic As (IA) and MMA/dimethylarsinic acid (DMA) ratios were used as indicators of arsenic metabolism. AS3MT 12590 genotype and HT 2 and 4 groups were significantly higher in arsenic exposed and in arsenic exposed group. Other SNPs and the role of other hand, MMA/DMA was explained by AS3MT 07395, 08979, 12590, and 14458 genotypes. Because association of AS3MT 14458 with arsenic metabolism observed in this study was consistent with other populations in other previous studies, this SNP may significantly affect the metabolism regardless ethnicity.

2300 Mechanism of Arsenic Carcinogenesis in Normal Human Lung Cells.

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Arsenic originates from both geochemical and numerous anthropogenic activities including mining, combustion of fossil fuels, wood preservation, agriculture and metallurgy. Exposure of the general public to significant levels of arsenic is widespread. Arsenic is a well-documented human carcinogen. Long-term exposure to low levels of arsenic in drinking water has been linked to bladder, lung, kidney, liver, prostate, and skin cancer. Among them, lung cancer is of great public concern. However, little is known about how arsenic causes lung cancer and few studies have considered effects in normal human lung cells. The purpose of this study was to determine the cytotoxicity and genotoxicity of arsenic in human primary bronchial fibroblast (NHFB) and epithelial cells (NHBE). Our data show that arsenic induces a concentration-dependent increase in cell death after short (24 h) or longer (120 h) exposures. Arsenic induces concentration-dependent but not time-dependent increase in chromosome damage in fibroblasts. No chromosome damage is induced after either 24 or 120 h arsenic exposure in epithelial cells. Using comet assay and gamma-H2A.X foci forming assay, we found that 24 h or 120 h exposure to arsenic induces increases in DNA double strand breaks in both cell lines. These data indicate that arsenic is cytotoxic and genotoxic to human lung primary cells. However, the mechanism of arsenic-induced genotoxicity could be different in bronchial epithelial cells than that in fibroblasts. This work is supported by NIEHS grant R15ES021587 (H.X.) and by NIEHS grant ES016893 (J.P.W.). The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.
2302 Changes in Regulation of Lipid Metabolism from Low-Dose Arsenic Exposure. 
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Arsenic (As) is naturally present in the environment, and it can be found at various levels in drinking water resulting from contamination of ground water and soil. As is a major health concern because it is associated with an increased risk of various diseases such as type II diabetes, cardiovascular disease (CVD), several types of cancer, and reproductive and developmental problems. Type II diabetes and CVD are in turn associated with altered blood lipid levels and obesity. Indeed, excess adipose tissue and altered blood lipid levels have been shown to be strong predictors for development of type II diabetes and CVD. Previous cell culture studies have shown that arsenic affects adipogenesis, and a recent in vivo study found that arsenic exposed dams displayed alterations in overall lipid metabolism and triglyceride levels. In the current study, we developed a model to investigate changes in transcription factors involved in regulating lipid metabolism. We exposed adult male C57Bl/6j mice to 0, 10, or 100 ppb As in drinking water for 6 weeks, and then isolated various tissues from these mice in the fed or fasting state. We analyzed expression of genes and proteins involved in lipid metabolism and regulation in adipose tissue and liver using quantitative PCR and western blotting. Our results show alterations in expression of transcription factor SREBP-1c and its target genes, including diacylglycerol acyltransferase (DGAT2) and fatty acid synthase (FAS). These results indicate that arsenic can alter the expression of genes and proteins involved in fatty acid synthesis and triglyceride production, which may contribute to how arsenic exposure may cause metabolic imbalances in exposed individuals. (Supported by NIH-IEHS Superfund Research Program (P42 ES007373), and NIH-NRSA Training Grant (2 T32 ES7272-21))

2303 Survey of Arsenic in Drinking Water in the Southern Gobi Region of Mongolia.
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Arsenic (As) is a naturally occurring toxicant of global concern. The extent of As content in ground water, however, has not been fully assessed in Mongolia, where all drinking water is sourced from groundwater. Mongolia is currently experiencing rapid mining development, especially in the southern region, which is triggering population growth and will drive increasing water demand for sale drinking water. Moreover, the high prevalence of As exposure and As related diseases just across the border in northern China further highlights the necessity for relevant studies in Mongolia. Thus, this study attempts to determine As concentrations in water sources near the Oyu Tolgoi mine in the Southern Gobi region of Mongolia and investigate its relationship with area, type and depth of the well, and develop a geographical map describing the spatial pattern of As concentration in the drinking water sources. The results of our study show that the current and the potential future exposure to As in the Southern Gobi region of Mongolia is significant. In terms of current exposures, almost half of the Herder’s wells that are currently in use and 16.4% of the Monitoring boresholes contain As levels above the World Health Organization’s recommended level. The results also indicate different levels of As concentrations in the water from different types of tube-wells even in the same area and a decreasing tendency of As concentration with increasing well depth suggesting different flow regions. As in all the drinking water sources in the Southern Gobi region of Mongolia is a critical public health issue, especially given the current mining boom in this region.

2304 The Role of miRNA-29B in Dysregulation of Mesenchymal Stem Cell Differentiation to Adipocytes by Low-Dose Arsenic Exposure.
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Human exposure to environmental toxicants is a well-known cause of disease and low chronic exposure may contribute significantly to longitudinal risk of chronic diseases. Pathogenic mechanisms for many environmental toxicants remain poorly defined, which limits development of effective interventions to protect against environmentally-derived chronic diseases. Low dose exposure to trivalent arsenic (AsIII) in drinking water is a major public health concern that contributes to a number of diseases and pathologies, including cardiovascular and metabolic diseases. While progress has been made in the understanding of the pathogenic signaling events contributing to arsenic-induced disease, many responsible mechanisms have not been elucidated. Control of miRNA expression and action presents a promising new means for understanding downstream effects of arsenic exposure; however, there are few reports of how arsenic regulates expression of miRNA and impacts their function. Our preliminary data indicated induction of miR-29 in white and brown adipose tissue isolated from arsenic exposed (100 μg/L in drinking water for 2 wk) mice and in human adipose-derived mesenchymal stem cells (hMSC) as arsenic inhibited adipocyte differentiation. Further analysis via real-time PCR revealed a 2-3 fold induction of miR-29b following arsenic exposure in hMSCs. The miR-29 family has been identified by multiple studies to be involved in cardiovascular and metabolic diseases, with reduced stem or progenitor cell differentiation capacity believed to be a fundamental means for disease progression. Analysis of downstream proteins by western blot indicates that C/EBP-zeta, an inhibitor of adipogenesis, is upregulated in arsenic treated cells and this upregulation is diminished in cells stably expressing an inhibitor of miR-29b. These data show that low-dose arsenic effects expression of miRNA in hMSCs, negatively affecting their ability to properly differentiate into adipocytes. Supported by NIHES grant R01ES013781.

2305 Human Exposure to Arsenic in Drinking Water Is Associated with Increased Protein Halogenation in Blood and Sputum.
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Chronic arsenic exposure to environmental high levels (> 200 ppb) in geologically contaminated drinking water has been correlated with increased incidence of chronic lung disease, including cough, bronchitis, shortness of breath and obstructive or restrictive lung disease. However, the effects of chronic arsenic exposures at lower levels (e.g., < 100 ppb), which are prevalent in the United States and near the current EPA MCL of 10 ppb, have not been well studied. Eosinophils and neutrophils are associated with lung disease, and these granulocytes selectively secrete peroxidases that halogenate protein tyrosines. A custom ELISA microarray platform was used to measure the halogenation of 24 individual proteins in paired sputum and serum samples from 55 subjects in 4 Mexican cities with differing arsenic levels. Eighteen halogenated proteins were significantly correlated (Spearman’s rank correlation, p<0.05) between the two fluids (highest r value =0.58). Halogenated protein results were compared to total arsenic levels in urine using multiple linear regression, accounting for the study participants’ age, BMI and resident city. The total urine arsenic was significantly associated (P ≤ 0.05) with the halogenation of 7 proteins in both plasma and serum (MMPI1, 2 and 9, EGF, VEGF, HBEGF, TGFIα) and 11 additional sputum-only proteins (PDGF, E-selectin, EGF, SP-A, leptin, HGF, CD14, IGF1, RANTES, TNF; ceruloplasmin) proteins. These results support the conclusion that environmental arsenic exposure is associated with elevated markers of pulmonary granulocyte activity. Replication in an independent study population is warranted to confirm our results. Supported by P42 ES4940, P30 ES006694, P50 CA057460, and U54 ES016015.
Rationale & Objectives: Risk due to dietary arsenic (As) exposure is specification dependent. For example, although elevated As concentrations are commonly observed in fish, shellfish, and seaweed, the health risks are generally assumed to be low due to the predominance of organoarsenicals (e.g. arsenobetaine). However, little specification data is available to support this assumption for many types of seafood. The objective of the research described in this poster is to fill this data gap by quantifying the relative contribution of various As species for a variety of store-bought seafood samples.

Methodology: Seafood samples were purchased from several supermarkets for As specification analysis using Liquid Chromatography Inductively Coupled Plasma Mass Spectrometry (LC-ICP-MS). The samples included cherry stone clams, oysters, shell clams, tiger prawn, halibut, mackerel, ling cod, marlin, snapper, yellow grouper, and Porphyra seaweed. Extraction of As species was performed with a methanol-ammonium carbonate solution using a DigiPREP block digestion system. Quantified As species included As(III), As(V), arsenobetaine, arsenocholine, dimethylarsinic acid, and monomethylarsonic acid.

Results: Total As concentrations ranged between 6 g kg⁻¹ (king mackerel) and 135 g kg⁻¹ (ling cod). The main contributor to As concentrations tended to be arsenobetaine, which ranged between 35% (yellow grouper) and 100% (king mackerel). As(V) was not a significant contributor (<1%) to total As concentrations in any of the samples.

Conclusions: Arsenobetaine was the major As species in most of the seafood samples tested. Future work will be done to assess whether As specification in foods is modified when digested in simulated gastrointestinal fluids.
Arsenic (As(III))-contaminated drinking water is a global health concern as chronic As(III) exposure poses risk for a number of cancers, diseases, and disabilities. As(III) exposure is associated with skeletal muscle weakness, impaired gait, and fatigue. While As(III) and other metals impact embryonic stem cell development, it is not clear how As(III) impairs adult stem cell functions in tissue maintenance and regeneration. We hypothesized that As(III) exposure affects adult skeletal muscle stem cell (MuSC) metabolism and function to disrupt muscle maintenance and repair capacity. To investigate this hypothesis, we examined muscle integrity, MuSC metabolism, and phenotype in hind limb muscles isolated from mice exposed to 0 or 100 μM As(III) for 5 weeks. Histological analysis demonstrated disrupted muscle bundles with peripheral fatty inclusions and ultrastructural analysis revealed large, fused muscle cell mitochondria in As(III) exposed mice relative to controls. MuSC isolated from the As(III) mice and cultured without As(III) for multiple doublings retained a phenotype with mitochondrial myopathy, autophagy, uncoupled oxidative phosphorylation, and impaired differentiation. This phenotype was a maladaptation to stress, as As(III) altered growth kinetics and resistance to oxidative stress. The phenotype and altered growth kinetics were reproduced in primary human MuSCs. These findings suggest that direct effects on MuSC mitochondrial phenotype, metabolism, and differentiation underlies muscle impairment observed following As(III) exposure. Supported by NIEHS grant R01 ES0136781, NIH K12 for Physical and Occupational Therapists (K12 HD055931) and NIA grant K01 1K01AG039477.

Bladder cancer has been associated with chronic arsenic exposure. Monomethylarsonic acid (MMAIII) is a metabolite of inorganic arsenic and has been shown to transform a human urothelial cell line (UROtsa). It was used as a model arsenical to examine the mechanisms of arsenical-induced malignant transformation of urothelium. A microarray analysis was performed to assess the transcriptional changes in UROtsa from chronic 50 nM MMAIII exposure that leads to transformation at three months. The analysis revealed only minor changes in gene expression at one and two months of exposure, contrasting with substantial changes observed at three months of exposure. To assess the changes occurring between 2 and 3 months of exposure, incremental analysis was performed for 29 genes, covering 7 distinct pathways (mitogenic, PI3K/AKT, apoptosis, JAK/STAT, oxidative stress, DNA repair, and inflammation), that were found changed at 3 months of expression based on the gene array analysis. Between 2 and 3 months of exposure, progressive alterations in the expression of several genes (i.e., PDGFRA, COX2, XAF1) were observed. These alterations are being correlated with expected phenotypic changes (i.e., hyper-proliferation, colony formation in soft agar) in the transforming cells. Since short-term exposure (up to two months) has not been shown to induce transformation, the gene expression changes observed for cultures treated up to 3 months and beyond suggest that a stress-threshold exists. This study was supported by the Superfund Basic Research Program Grant (NIH grant ES04940) from National Institute of Environmental Health Sciences, and the Taube in Toxicology and Toxicogenomics (NIEHS grant ES05609). Additional support from NIH grant CA23074, and NIEHS grant ES06694.

Arsenic trioxide is established as one of most effective drugs for treatment of patients with acute promyelocytic leukemia (APL). However, non-promylocytic leukemias HL-60 cells exhibit resistance to arsenic. An improved understanding of the underlying resistance mechanism for As2O3 and its biomethylation products, namely, monomethylarsonic acid (MMAIII) on the treatment of tumors. In the present study, we investigated the molecular mechanisms underlying iAsIII and its intermediate metabolite MMAIII-induced antitumor effects in the HL-60 cells. Here, we show that the HL-60 cells (MuSCs) exhibit resistance to inorganic arsenite (IC50=10 μM), but are relatively sensitive to its intermediate MMAIII (IC50=3.5 μM). Moreover, we found that the multidrug resistance protein 1 (MRP1), but not MRP2, are expressed in HL-60 cells, which reduced the intracellular arsenic accumulation, and conferred resistance to inorganic iAsIII and MMAIII. Pretreatment of HL-60 with MK571, an inhibitor of MRP1, significantly increased iAsIII and MMAIII-induced cytotoxicity and arsenic acid levels, suggesting that the expression of MRP1/4 may lead to HL-60 cells resistance to trivalent arsenic compounds.

Diabetes mellitus is a metabolic syndrome characterized by inappropriate production of insulin or the inability of cells to respond to insulin. It is estimated that by the year 2050, 1 in 3 U.S. adults will have diabetes mellitus. Insulin is the principal hormone involved in lowering blood glucose and functions by suppressing liver glucoseogenesis and glycogenolysis and by stimulating the glucose uptake in skeletal muscle and adipocytes. Recent epidemiological studies both in the USA and abroad have linked chronic ingestion of low levels of inorganic arsenic (iAs), an environmental toxicant, to the onset of diabetes mellitus. Although these observations have been met with some skepticism, there are few mechanistic studies to elucidate the mechanisms by which iAs perturbs insulin signaling. The few studies that have been performed have focused namely on adipocyte in-vitro models. Here we show that L6 myocytes, an insulin responsive cell line, exposed to low doses of iAs (0.25 to 2 μM) for 4 or 7 days showed a decreased insulin stimulated glucose uptake but no decrease in phospho-AKT or phospho-AS160. In addition, we found that phospho-ERK signaling decreased, while phospho-p38 MAPK signaling increased in response to prolonged iAs treatment. Intriguingly enough increased p38 MAPK activity has been associated with insulin resistance. These data support the epidemiological evidence that chronic exposure to low, physiologically relevant levels of arsenite can contribute to insulin resistance and type 2 diabetes, and that the mechanisms involved are different than those produced by iAs in adipocyte cell models. While and the etiology of type 2 diabetes has yet to be elucidated these data show that in addition to pharmacological treatment and lifestyle modifications, environmental exposures should also be considered when evaluating the etiology of type 2 diabetes.
An integrative evaluation of the toxicopathological effects of aflatoxin B1 (AFB1) was conducted in this study. Briefly, male F344 rats were orally exposed to a single-dose of AFB1 at 0, 50, 250 or 1000 μg/kg body weight (BW) or repeatedly dosed of AFB1 at 0, 5, 10, 25 or 75 μg/kg BW for up to 5 weeks. Biochemical and histological changes were assessed together with the formation of AFB1-lysine adduct (AFB-Lys) in serum and liver foci positive for placental form glutathione S transferase (GST-P+). In single-dose protocol, serum ALT, AST and AFB1 were dose-dependent elevated with maximal changes (> 100 fold) in AFB1 at 3-day after treatment. Animal that received 250 μg/kg AFB1 showed concurrent bide duct proliferation, necrosis and appearance of GST-P+ hepatocytes at 3-day while the pre-neoplastic GST-P+ foci appeared after 1-week. Neither liver GST-P+ hepatocytes nor foci were induced by 50 μg/kg AFB1 treatment. In repeated-dose protocol, bide duct proliferation and liver GST-P+ foci co-occurred after 3-week, followed by proliferation of foci formation after 4-week and dramatic ALT, AST and CK elevations after 5-week exposure in animals received 75 μg/kg AFB1. Liver GST-P+ hepatocytes and foci appeared in a dose- and time-dependent manner, low dose of AFB1 after 5-week exposure in animals received 75 μg/kg AFB1. Liver GST-P+ hepatocytes and foci were assessed together with the formation of AFB1-lysine adduct (AFB-Lys) in serum and liver foci positive for placental form glutathione S transferase (GST-P+) in serum. ALT, AST and CK elevations further assessed with various demographic parameters. These results demonstrated the temporal patterns of AF exposure in rural human populations of Uganda.

In this study, two experiments (in vivo and in vitro) were carried out to study the effects on OTA and FB1 on pigs mononuclear cells as well as on live pigs to assess the effects of these mycotoxins on their health with particular reference to the immune system and pathomorphological changes. The in vivo study showed a time vs concentration decrease of pig mononuclear cells were used the MTT assay and exposed to 5 and 40 ng/ml for ochratoxin A and 5 and 40 μg/ml of FB1 at 12, 24 and 48 hrs. While the in vivo study showed the ochratoxin A induced hepatic cell viability decrease as compared to FB1 when exposed singularly, whereas exposure to both mycotoxins simultaneously showed further reduction of cell viability. In the in vitro study was done with mycotoxic nephropathy induced in eighteen young pigs with mouldy diets containing 0.5 ppm ochratoxin A (OTA) and/or 10 ppm fumonisin B1 (FB1) for three months. Exposed porcine damage provoked by OTA was seen in the kidneys, as expressed by the strong degenerative changes in proximal tubules and fibrosis in kidneys, FB1 was found to induce an increase in permeability of vessels mainly in lung, brain, cerebellum or kidneys and slight to moderate degenerative changes in kidneys. The exposure to both mycotoxins simultaneously revealed synergistic pathomorphological changes characterized by the combination of the main lesions provoked by each mycotoxin alone, being stronger in their expression when administered together. In addition, Exposure to both mycotoxins and their combination showed induced humoral immune response in all experimental pigs shown by decrease in antibody titers.
DON-induced CCK release in STC-1 cells. The results suggest that DON might induce CCK-release in enterodocrine cells by increasing CaSR sensitivity to extracellular [Ca2+]0 which mediates increased [Ca2+]i influx via L-type-VSCCs. DON-induced calcium and hormonal responses demonstrated herein might ultimately contribute to anorexia and emesis potentially as a protective mechanism following ingestion of the toxin.

PS 2321 Red Clover Exhibits Multifaceted Activity on Breast Cancer Cells.

An increased breast cancer risk for postmenopausal women taking hormone therapy has been demonstrated by the Women's Health Initiative. Botanical dietary supplements, including red clover products, are commonly used to alleviate menopausal symptoms but still lack efficacy and safety studies. Red clover isoflavones, such as biochanin A and genistein, have been promoted for their cancer preventive activity, in part due to genistein's ERβ selective activity. Other studies suggested possible side effects in estrogen sensitive tissue due to genistein's ERα mediated estrogenicity. This study examines these controversial claims by performing several estrogenic assays with a well-characterized clinical red clover extract and pure isoflavones. Proliferation experiments in ERα breast cancer cells (MCF-7) showed that the effect of the red clover extract on cell proliferation is highly concentration dependent and has a bell-shaped curve: Low- and high concentrations (30 - 100 ng/mL and ≥ 5 μg/mL, respectively) reduced cell proliferation, whereas intermediate amounts (300 ng/mL - 1 μg/mL) increased it. The pure isoflavones, genistein and biochanin A, exhibited similar bell-shaped dose dependencies. Low-(10 nM) and high isoflavone concentrations (> 10 nM) decreased cell proliferation, whereas cell survival increased at 300 nM - 1 μM. These findings are in agreement with prior reports of genistein preferentially binding to ERβ at ~6 - 20 nM and to ERα at ~300 nM, which may explain the dose-dependent effects of the isoflavones and the extract. Supporting the proliferation results, ERE-luciferase and gene induction assays showed that red clover extract at 300 ng/mL - 1 μg/mL mimics estradiol and can potentially have similar adverse effects if taken in high quantities. These findings highlight the importance of correct dosing for isoflavone-containing products and the relevance of dose considerations to ensure safety of botanical dietary supplements.

PS 2322 Occurrence of T-2 Toxin, HT-2 Toxin and Zearalenone in Retail Foods in Japan.
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Fusarium toxins are a group of mycotoxins produced by many kinds of Fusarium species, and frequently detected in field crops such as wheat, barley, maize and corn. They cause mycotoxicosis, a serious health hazard to humans and domestic animals. Therefore, collecting the information about fusarium toxin contamination in daily foods is crucial. Occurrence data of deoxynivalenol, a major fusarium toxin, has been collected in worldwide, but that of other toxins is limited. In this study, we examined foods from Japanese retail shops for contamination with three kinds of fusarium toxins, T-2 toxin, HT-2 toxin and zearalenone. Food samples were extracted with methanol-water (75 : 25). After filtration, the extract was diluted five times with phosphate buffered saline, and subjected to a DZT Bioavailability of Aflatoxin, Fumonisin and Aflatoxin/Fumonisin Mixtures in Fischer 344 Rats.

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Aflatoxins (AFs) have been linked to hepatocellular carcinoma and mortality in humans and animals. Chronic AF exposure, particularly in developing countries, is a significant problem that continues to contribute to public health issues in communities burdened by poor economic status and food scarcity. One possible strategy to...
improve food safety is to reduce exposures to AF using montmorillonite clay which binds AFs in the GI tract and decreases bioavailability of these toxins. Our recent work in Ghana has shown that communities at risk for AF exposure are also at high risk for fumonisin (FB) exposure. In this study, montmorillonite clay was tested for FB binding capacity in combination with AFB, in a rodent model. Fisher-344 rats were gavaged once with 0.125 mg AFB/kg b.w. and/or 25 mg FB/kg b.w. following an acclimation period with feed containing clay additive. Urine samples were collected at 12 hr time intervals for 72 hr following gavage and were analyzed for AF and FB biomarkers. Lower AFM1 and FB1 levels in the urine were indicative of reduced absorption of the parent compounds through the GI tract and into the circulation. AFM1 and FB1 excretion peaked between 12-24 hr and quickly declined by 36 hr. Addition of clay at 0.25%, 0.5% and 2% w/w feed decreased AFM1, excretion by 88-97% at the 12 hr time point, indicating highly effective binding. Clay treated animals also had a reduction in FB1 excretion, but to a lesser extent than the AFM, biomarker (i.e. 41-80%). When in combination both AFM1 and FB1 binding occurred, but capacity was decreased by almost half, suggesting that AFB and FB are competing for similar binding sites on the clay. This study indicates that inclusion of montmorillonite clay in contaminated diets can reduce bioavailability of both AFs and FBs in populations at high risk for mycotoxin exposure. (Supported by NIH/NCRMHD RO1-MD00519-01.)

2326 Development and Validation of an Analytical Method for Vomitoxin in Gavage Dose Formulations Used in Rodent Toxicology Studies.
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Vomitoxin, also known as deoxynivalenol (DON), is a trichothecene mycotoxin produced by certain types of Fusarium fungi. It occurs predominantly in grains such as corn, wheat, barley, and rice, and has been shown to have great stability during storage, processing, and cooking of food. Because of the potential for widespread contamination of food and exposure to humans through ingestion, the National Toxicology Program is investigating the toxicity of vomitoxin.

Thus, the current study was undertaken to develop and validate a formulation analysis method for vomitoxin in deionized water as a gavage vehicle for use in toxicology studies. Important objectives of an analytical method for supporting a toxicological study are to insure that the correct test article is being administered at the specified dose concentrations and that the dose formulations are stable. To achieve these objectives, an Ultra Performance Liquid Chromatography (UPLC) method was developed and validated for analysis of vomitoxin in deionized water. Sample preparation involves a simple dilution with water and addition of an internal standard solution (2,6-dimethylphenol in acetonitrile). The method was successfully validated over the range 1.5 to 720 μg/mL and the limit of quantitation was estimated as 0.8 μg/mL. Assessment of formulation stability at ambient and refrigerated storage conditions, over a 42-day period was evaluated. Dose simulation stability testing was also conducted to evaluate the formulation stability for a period of up to three hours under simulated dosing conditions. In addition, analysis period stability of the analytical preparations was assessed. This method will be used to support toxicology studies of vomitoxin conducted by the National Toxicology Program.

2327 Comparative Hepatic Glutathione S-Transferase Mediated Detoxification of Aflatoxin B1 in Chickens.
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Efficiency of hepatic glutathione S-transferase (GST)-mediated conjugation of bioactivated aflatoxin B1 (AFB1) is critical to species resistance to this toxic and carcinogenic mycotoxin. Poultry are among the most susceptible animals, and domestic turkeys are especially susceptible, a condition we have shown to be associated with a deficiency of AFB1-detoxifying GSTs. Chickens are more resistant than domestic turkeys, yet little is known about their hepatic GST detoxification capabilities. In this study, we compared hepatic GST-mediated detoxification of prototype substrates 1-chloro-2,4-dinitrobenzene (CDNB), 1,2-dichloro-4-nitrobenzene (DCNB), ethacrynic acid (ECA), and cumene hydroperoxide (CHP) and toward the exo-AFB1-8,9-epoxide (AFBO) in livers of Broiler, Fayoumi, and Leghorn chickens. All breeds had equivalent GST activities, except Fayoumi, which was significantly lower toward CDNB. Broiler, Fayoumi, and Leghorn had similar GST-mediated conjugation activities toward AFBO (6.5 ± 0.2, 6.1 ± 0.6, 6.4 ± 0.2 pmol/min/mg, respectively). This level of hepatic AFB1 detoxification is substantially higher than that reported for in vitro conjugation of AFB1 by GST, which is an accurate predictive marker species resistance to this mycotoxin. Supposed in part by NRI Competitive grant 2007-35205-17880 from the USDA-NRI Animal Genome Project.

2328 Bisphenol A (BPA) Levels in Liquid Supernatants of Canned Foods Determined by Highly Sensitive BPA ELISA.
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BPA [2,2-(4,4'-dihydroxydiphenyl)propane], an endocrine disruptor, mimics the action of 17beta-estradiol (E2) in mammals, increasing the risk of hormone-related health problems such as early puberty, infertility, breast, ovarian and prostate cancers and insulin resistance. Fetuses and newborns are most vulnerable to the BPA toxicity. Recently, BPA levels of canned soup solids were measured by LC/MS/MS (detection limit, 2 ng/g) and it was found that the soup solids contained 10 to 80 ng/g BPA. A subsequent study revealed that the group that consumed a 12-ounce serving/day, for 5 days of canned soups, excreted 19-fold higher BPA in urine (mean, 21 ng/ml) compared with the control group that consumed similar amounts of fresh soups (mean, 1.1 ng/ml). BPA in liquid supernatants of canned foods contain -10-fold lower levels of BPA compared to the solids. To screen BPA in supernatants, a highly sensitive and facile BPA ELISA has been developed after production of BPA polyclonal antibodies by immunization of a goat with carboxyalkyl-derivatized BPA conjugated to KLH. The detection limit of the BPA ELISA was 1 pg. Whereas anti-BPA slightly cross-react with BPS, BPF or resveratrol, which are structurally similar to BPA. Ten-fold diluted supernatants of canned soups were applied to BPA ELISA. Supernatants obtained from 3 kinds of soups (3 cans/each kind of soup, 9 data points) produced by first company contained 9.56 ± 0.96 ng/ml, 9.07 ± 0.38 ng/ml and 10.38 ± 0.83 ng/ml of BPA, similar to ~8.70 ng/ml of BPA levels in supernatants of second company. A negative control, supernatants of canned vegetable soup, contained extremely low levels of BPA (0.05 ± 0.02 ng/ml), suggesting use of BPA-free can linings. These results demonstrate that the competitive BPA ELISA is suitable for measurements of BPA leaching from the epoxy film-coated cans using liquid supernatants from canned foods.

2329 Perinatal BPA Exposure at Low Doses Impairs Oral Tolerance and Immunization to Ovalbumin in Offspring Rats at Adulthood.

Aims: Bisphenol A (BPA) used in food packaging impacts gut epithelial barrier after perinatal exposure. Because oral desensitizing by gut epitopes is a major mucosal immune response, our aim was to address the consequences of perinatal BPA exposure at low doses on oral tolerance and immunization at adulthood. Methods: Dams were treated per os from gestation day 15 to pup weaning with BPA [0.5, 5 or 50 μg/kg/d] or vehicle (corn oil). Female offsprings (day 45) were used to assess paracrine and transcellular jejunal permeability by Using chambers, and immune response to ovalbumin (OVA) after either oral tolerance, immunization or oral challenge. Results: Perinatal BPA exposure decreased jejunal paracellular permeability by 2-fold at 0.5 and 50 μg/kg/d, and by 3-fold at all doses for transcellular permeability (p<0.05). BPA exposure at 5 and 50 μg/kg/d increased anti-OVA IgG1 titers after an oral tolerance protocol (116±62x103 and 86±29x103 respectively vs 7±2x2 ±103 in controls; p<0.04). Anti-OVA IgG1 titers were only increased at 5μg/kg/d after OVA immunization. Enhanced humoral response in rats exposed to BPA [0.5, 5 or 50 μg/kg/d] was associated with higher IFN-γ concentration (2-fold) and MLN of OVA-tolerized (20-fold) rats (p<0.05). Final, oral OVA challenge in BPA group increased MPO activity (316±15x103 in controls; p<0.04) and decreased TGF-β concentration (2-fold, p<0.05). IFN-γ concentration (2-fold, p<0.05) and decreased TGF-β concentration (3-fold, p<0.05) in the colon, indicating inflammation. Conclusion: Perinatal exposure to low doses of BPA decreased jejunal paracellular and transcellular permeability, and impaired oral tolerance and immunization to dietary antigens at adulthood. BPA treatment during perinatal period affects intestinal homeostasis in a nonlinear dose-response relationship. These results suggest that perinatal period is a critical window for BPA exposure that may trigger food intolerance in later life.
2330 Perinatal BPA Exposure at Low Doses Impairs Immune Homeostasis and Promotes Intestinal Parasite Infection in Young Rats.


Aims: Perinatal exposure to 5µg/kg/day of the food contaminant bisphenol A (BPA) impaired oral tolerance in adult rats (see Méndar et al, abstract 1). Herein, we aimed to address the consequences of BPA perinatal exposure on immune homeostasis in young rats at weaning, i.e. oral tolerance, systemic immunization with ovalbumin (OVA) and parasitic infection. Methods: Dams were given per os from gestational day 15 to pup weaning BPA [5µg/kg/d] or vehicle (corn oil). Weaned female offspring (day 25) were used to assess para- and trans-cellular jejunal permeability by Using chambers, and immune responses following oral toler- ance or immunization to OVA, or after infection with a gut nematode Nippostrongylus brasiliensis (N brasil). Results: Perinatal treatment with BPA did not affect intestinal permeability or humoral response (anti-OVA IgG titers) following either oral tolerance or immunization in D25 rats. However, a decrease of OVA-induced IFNγ secretion was observed in spleen of OVA-sensitized rats (53±15 vs 317±162 pg/ml; p<0.05) and in mesenteric lymph nodes of OVA-toler- ized rats (3.9±2.3 vs 19±18 pg/ml; p<0.05). The lack of cellular response to food antigens questioned the ability of BPA-exposed rats to clear intestinal infections. A 3-fold increase in N brasil living larvae was observed in the intestine of BPA-exposed rats compared to controls (2817±689 vs 757±291 larvae/g of faces, respectevly; p<0.05), but no significant change in myeloperoxidase activity into jejunal tissues. Conclusion: Perinatal exposure to low dose of BPA did not affect humoral response to the food antigen OVA in juvenile rats. However, a decrease of OVA-induced IFNγ secretion in BPA-exposed rats emphasized a lack of specific cellular response to food antigens. Finally, perinatal BPA treatment evoked an increased risk to intesti- nal parasitic infection without triggering an inflammatory response, demonstrating impaired immune defence in early life stages.

2331 Perinatal Periportal Exposure to Bisphenol-A Increases Hepatic Steatosis in Immature and Adult Mice.

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While diet and physical activity remain the predominant reason for development of obesity and obesity-related disease, there is some concern that environmental expo- sure to chemicals may be a predisposing factor. Bisphenol B (BPA), a component used in the manufacturing of certain plastics and plastic resins, can leach into food and/or drink from food and beverage containers. High urinary BPA levels have been positively associated with general and abdominal obesity in human popula- tions. In rodents, developmental BPA exposure increases body weight and fatty liver (steatosis). The purpose of this study was to uncover potential mechanisms by which perinatal-periportal (PNPP) exposure to BPA increases liver steatosis. Pregnant CD-1 mice were administered 25 or 250 µg BPA/kg/day (BPA25, BPA250, respectively) via osetric pump, and then after weaning on PND 20, the resulting daughters were exposed to BPA via drinking water up through PND 35. Tissues were collected at PND32 and at 39 weeks of age. Livers were analyzed for triglyceride content, stained with Oil Red O and relative mRNA expression was quantified by qPCR. At PND32, BPA25 and 250 increased liver Oil Red O staining compared to controls. BPA25, but not BPA250, increased protein expression for lipogenic enzymes, (Acc-1 and Fas) but did not significantly increase lipogenic gene expression. In adult mice, BPA25, but not BPA250, increased Oil Red O staining. BPA25 increased Acc-1 expression, whereas BPA250 increased Srebpl-c, Acc-1, and Fas. Protein expression was not changed. In conclusion, PNPP exposure to BPA had some effect in promoting steatosis and pro-steatotic gene expression in immature and adult female mice, but the exact mechanism by which BPA promotes steatosis remains unclear.

2332 Leaching of Chemicals with Estrogenic Activity from BPA-Free Materials after Common Use Stresses.

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Since 2007, many consumer products are no longer made from polycarbonate (PC) plastics because of widespread concern about the estrogenic activity (EA) of bisphe- nol-A (BPA). However, BPA-Free plastics have not been thoroughly vetted by most product manufacturers to ensure that they do not release other chemicals that have EA. We have used a robotized MCF-7 and BGI Luc assays currently undergoing val- idation by ICCVAM/NICEATM to quantify the total EA in chemicals leaching from a variety of widely-available plastic resins used to make consumer products and packaging. This study expands the prior work of Yang et al, 2011(EHP 119:989- 998). We assayed the total EA in chemical mixtures leaching from unstressed and stressed polycarbonate (PC), polypropylene (PP), polyethylene (PE), amine olefin copolymer(COC), and polyethylene terephthalate glycol-modified (PETG - Tritan™) plastic resins. Plastic resins were subjected to simulated common-use stresses of dishwashing, microwaving, and sunlight, and then extracted for 4-72 hours at 73 degrees Celsius by saline (hydrophilic) or ethanol (hydrophobic) sol- vents. The total EA in these leachates were then quantified and EA-specific re- sponses were validated by conducting confirmation assays. EA positive determinations were made on samples with values 3 standard deviations or more higher than vehicle controls. Using these criteria, Tritan PETG and PC resins were consistently significantly positive for EA. PE and PP samples often tested positive for EA, but some consistently had no detectable EA. COC resins had no detectable EA. Our data show that BPA-Free often does not mean EA-Free, i.e., that leaching of chemicals having EA from the plastic resins is often not addressed by simply choosing BPA-Free materials.

2333 Microsomal Metabolism of Asarone Isomers.

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The alkenylbenzenes alpha-asarone (αA; CAS 2883-98-9), beta-asarone (βA; CAS 5273-86-9), and gamma-asarone (γA; CAS 5353-15-1) are constituents of various plants, e.g. Acorus calamus (Sweet Flag) and some peppers. Both, αA and βA are carmogenic to rodents and exhibit genotoxic effects in vitro. Neither genotoxicity nor carcinogenicity of γA have been evaluated so far. Several alilic alkenylbenzenes such as safrole, estragole and methyleugenol are well known genotoxic carcinogens, whereas their proenolypic analogues are not. This suggests that an allylic side chain, and therefore the capability for the formation of an αOH metabolite, may be a basic prerequisite for the carcinogenicity of those compounds. However, the propenyl compounds αA and βA are an exception of that “allylic rule”. We suggest the ortho-methoxy groups of αA and βA to be a key structural element for the mode of action of their carcinogenicity.

We investigated the metabolism of αA, βA and γA using liver microsomes from dif- ferent species (including human). Identity of metabolites was confirmed by LC-MS/MS and 1H-NMR spectroscopy in comparison with synthesized reference standards.

Our results show that the side chain hydroxylation of αA and γA was the predomi- nating metabolic step leading to E-3’OHA (from αA) and Z-3’OHA (from γA), respectevly, together with the formation of side-chain dihydrodiols, for which the epoxides may be the precursors. To a smaller extent, we found enzymatically formed secondary metabolites derived from the alcohols like 3’oxoα and 1’oxoα, but no corresponding carboxylic acids. Furthermore, we found the corresponding mono-demethyliated phenolic metabolites. These results are comparable to our previous results on phase I metabolism of methyleugenol and methyleugenol. In contrast, βA showed a more complex pattern of metabolites: in addition to Z- 3’OHA, the unexpected direct formation of 1’OHA was proven. This metabolic step, possibly facilitated by steric attraction of the Z-configurated βA side chain and the ortho-methoxy substituent, may explain the carcinogenicity of βA, but not of αA, where no 1’OHA metabolite was found.

2334 Histamine in Scombrotxin Fish Poisoning: Toxicology, Epidemiology and Dose-Response Analysis.

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Histamine plays important physiological functions such as immune responses and gastric acid secretion. However, ingestion of fish containing large amounts of spoilage-originated histamine can result in scombrotxin fish poisoning (SFP), a common chemical-originated food poisoning that affects cardiovascular, gastroin- testinal and neurological systems. Fish importing countries have established regulations and limits for histamine in fish and fishery products to protect con- sumers. However, some limits were established in a pre-quantitative risk assessment era. An assessment of toxicological and epidemiological data indicates that though other biogenic amines might also play a role in the etiology of SFP, histamine is the most significant causative agent. Due to limitations of the disease-reporting system, the epidemiology-based dose-response approach is not suitable in developing a safety limit of histamine in fish. Instead, a dose-response analysis was conducted
based on human oral challenge studies selected. Both the NOAEL and BMD assessments identified 50 mg of histamine per meal as the dose where either adverse effects were not noted or the estimate of additional risk (lower confidence level) was low. At this level healthy adults would not be expected to exhibit any of the symptoms associated with SFP. This dosage level will not apply to children and individuals with a specific sensitivity to histamine. In addition, this level was derived from data on small number of subjects. While the variation of response appears to be reflected in the study results, further studies would be most helpful in refining this threshold value. Using a conservative serving size of 250 g fish meat/meal, the maximum concentration of histamine in fish that should not cause an adverse effect was calculated as 200 ppm. Compliance sampling plan can be derived based on this threshold level to ensure a desirable level of protection of the public health.

2335 Carrageenan-Induced Disruption of Mucosal Barrier via Regulation of Proinflammatory NF-κB and Early Growth Response Gene 1: A Mechanistic Implication of Food-Borne Inflammatory Bowel Disease.

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The widely used food additive carrageenan (CGN) has been shown to induce intestinal inflammation, ulcerative colitis-like symptoms, or neoplasm in the gut epithelium in animal models, which are also clinical features of human inflammatory bowel disease. NF-κB or EGR-1 aggravated barrier disruption by CGN, which is clinically evident in the formation of urinary MAs and of N7-Gua DNA adducts in liver, kidney and lung was measured 16 h after application using HPLC-MS/MS. At the lowest dosage (0.1 μg AA/kg bw), N7-Gua DNA adducts were below the limit of detection in any organ tested. At 1 μg/kg bw, enhanced adduct levels were found in kidney (-1 adduct/106 nucleotides) and lung (-<1 adduct/106 nucleotides), but not in liver. At 10 and 100 μg/kg bw, adducts were found in all three organs not significantly different to those found at 1 μg AA/kg bw (about 1-2 adducts/106 nucleotides). In two pilot intervention studies, MAs of AA and AC were monitored after ingestion of test meals of 150 g self-made potato chips (1 mg AA, 13 volunteers, study 1) and of 175 g commercially available potato chips (44 μg AA and 5 μg AC, 5 volunteers, study 2). Urinary MA contents were determined by HPLC-MS/MS for up to 72 h (study 1), respectively 24 h (study 2). Total excretion of AC-related MA exceeded that of AA-related MA by factors of about 12 (study 1) to 4 (study 2). These results suggest markedly higher exposure to AC than to AA from heat treated potato based foods representing a major source of dietary exposure to AA. They mandate further research to broaden the database on dietary AA and AC.

2336 Comparison of the Estrogenic Activity of Licorice Species with Hops in Botanical Dietary Supplement Formulations for Women’s Health.


The Women’s Health Initiative showed an increased risk of breast cancer for postmenopausal women taking hormone therapy. As a result, many women have turned to botanical supplements to manage menopausal symptoms, although there is limited data about their efficacy and safety. Our previous studies demonstrated estrogenic and chemopreventive properties for hops (Humulus lupulus). The goal of the current study was to compare the estrogenic effects of three common licorice species (Glycyrrhiza glabra, Glycyrrhiza uralensis, Glycyrrhiza inflata) with those of hops. Methanol extracts of the licorice species showed a dose-dependent induction of an estrogen responsive alkaline phosphatase in endometrial cancer cells with Glycyrrhiza uralensis being the most potent. Compared to hops, the activity of the licorice species was significant but less pronounced. Similar results were obtained in an estrogen response element (ERE)-luciferase reporter assay in breast cancer cells. The licorice constituent, liquiritigenin, was the major ligand of estrogen receptor (ER) in pulsed ultrafiltration mass spectrometry of the licorice extracts. Competitive binding assay using purified human ERs showed liquiritigenin as a selective ligand of ERβ. In comparison to 8-prenylnaringenin (8-PN), the major estrogenic principle of hops, liquiritigenin had lower affinity to both ERs. Liquiritigenin was less active than 8-PN in the induction of alkaline phosphatase and ERE-luciferase. Isoliquiritigenin, the precursor chalcone of liquiritigenin, demonstrated significant activity in these estrogenic assays, while xanthohumol, the precursor of 8-PN, did not exhibit estrogenic effects. The estrogenic activity of isoliquiritigenin was partially associated with its cyclization to liquiritigenin. These data suggest that licorice species are moderately estrogenic and are worthwhile further exploration as effective and safe dietary supplements to alleviate menopausal symptoms.

2337 Acrylamide and Acrrolein: Heat-Induced Contaminants in Food.

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The genotoxic carcinogen (IARC class 2A) acrylamide (AA) is a food contaminant formed by thermal treatment of food. This applies as well to acrolein (AC) an α,β-unsaturated aldehyde grouped into IARC group 3B. Whereas human dietary exposure levels to AA are well established and range between about 0.5-5 μg/kg bw/d, exposure to AC is much less investigated. After uptake, AA is partly metabolised into the genotoxic glycidamide (GA). GA forms DNA adducts, primarily at N7 of guanine (N7-GA-Gua). AA, GA and AC are conjugated to glutathione (GSH) and excreted via urine as mercapturic acids (MA). AA was given by gavage in single doses of 0.1-10,000 μg/kg bw to female rats. Formation of urinary MAs and of N7-GA-Gua DNA adducts in liver, kidney and lung was measured 16 h after application using HPLC-MS/MS. At the lowest dosage (0.1 μg AA/kg bw), N7-GA-Gua DNA adducts were below the limit of detection in any organ tested. At 1 μg/kg bw, enhanced adduct levels were found in kidney (-1 adduct/106 nucleotides) and lung (<1 adduct/106 nucleotides), but not in liver. At 10 and 100 μg/kg bw, adducts were found in all three organs not significantly different to those found at 1 μg AA/kg bw (about 1-2 adducts/106 nucleotides). In two pilot intervention studies, MAs of AA and AC were monitored after ingestion of test meals of 150 g self-made potato chips (1 mg AA, 13 volunteers, study 1) and of 175 g commercially available potato chips (44 μg AA and 5 μg AC, 5 volunteers, study 2). Urinary MA contents were determined by HPLC-MS/MS for up to 72 h (study 1), respectively 24 h (study 2). Total excretion of AC-related MA exceeded that of AA-related MA by factors of about 12 (study 1) to 4 (study 2). These results suggest markedly higher exposure to AC than to AA from heat treated potato based foods representing a major source of dietary exposure to AA. They mandate further research to broaden the database on dietary AA and AC.

2338 Sex Hormone Modulation of Long-Term Induction and Short-Term Inhibition of CYP1A1/2 by Genistein in HepG2/C3A Cells.

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Genistein is widely consumed in soy products and dietary supplements for its reported beneficial health effects including cancer prevention. However, there have been conflicting data suggesting that genistein ingestion has both anticancer and cancerc-promoting activities. Cytochromes P450 (CYP1A1) play key roles in the metabolic activation of many carcinogens. Since genistein has both anti-estrogenic and estrogenic activities, sex-specific factors could also contribute to its biological activities. In the current study, human hepatoma HepG2/C3A cells were cultured in media with defined human sex hormone profiles to investigate CYP1A1 inhibition and induction by genistein. In male hormone supplemented cells, CYP1A1 and CYP1A2 gene expression and activities were induced to higher extent compared with female hormone supplemented cells after either β-naphthoflavone or genistein treatment. However, basal gene expression of CYP1A1 and CYP1A2 were higher in female- than in male-hormone supplemented cells. In the inhibition studies, CYP1A1 activities were significantly lower in the male-specific medium than in female specific medium. The results showed that genistein could exert both long-term (3-day) induction and short-term (1-hr) inhibition effects on CYP1A1 activities in vitro which could explain the inconsistent cancer-related reports. Furthermore, there were significant differences in both the inductive and inhibitory effects of genistein between the male- and female-specific media suggesting that sex hormones, in physiological concentrations and ratios, can modulate the effects of genistein on CYP1A1 gene expression and activities in a human liver cell line.

2339 Safety Evaluation of Optimash BG from T. Reesei for Use in Grain, Brewing, and Carbohydrate Processing.

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Optimash BG is a cellulase enzyme used in grain processing (for the production of potable alcohol and brewing) and glucose production from starch. The enzyme, endo-glucanase 2, is produced from a recombinant modified strain of Trichoderma reesei. A battery of toxicology studies was conducted to investigate its potential to
cause adverse effects in humans using methods complying with OECD guidelines. All studies were conducted according to Good Laboratory Practice. Acutely, Optimash BG is not an eye irritant, a very mild skin irritant, and not toxic by ingestion with an oral LD50 greater than 2000 mg/kg bw. Optimash BG is not a mutagen, a clastogen, or an aneugen. In the in vitro cytogenetic test using cultured human lymphocytes cells, Optimash BG did not induce chromosomal aberrations (both structural and numerical) in the presence and absence of metabolic activation (S-9 mix) up to the highest concentration (5000 ug TP/ml). No mutagenic activity was noted in the Ames assay in the absence and presence of S-9 mix up to 5000 ug TP/plate. In a repeated 90 days oral (gavage) in Wistar rats, no biological or statistical differences were observed. Also, no treatment-related changes in the hematology and clinical chemistry were noted at study termination. The NOAEL was established at 80 mg total protein/kg bw/day (97.6 mg TOS/kg bw/day). Under the worst-case scenario that Optimash BG is applied at the maximum rate and the enzyme is neither destroyed nor removed during processing, the use of the enzyme in grain processing, carbohydrate processing and brewing is not expected to result in adverse effects to humans. With a margin of safety of 258 and a PADI of 39%, the use of Optimash BG is not of toxicological concern.

**2340 A 13-Week Subchronic Toxicity Study of Glycodil Fatty Acid Esters in F344 Rats.**


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Glycodil fatty acid esters (GEs) have been recently identified as food process contaminants in refined edible oils. Although there is toxicological concern arising from potential release of glycodil from parent esters during digestion in the gastrointestinal tract, little is known about in vivo toxicity of GEs. In the present study, subchronic toxicity of two types of GEs, oleate and linoleate esters, was investigated with administration at concentrations of 0, 225, 900 and 3600 ppm (equivalent molar concentration to 800 ppm glycodil) in drinking water for 13 weeks to male and female F344 rats. For comparison, treatment with 200 and 800 ppm of glycodil was also performed. Body weight gain of both sexes was markedly reduced with 800 ppm glycodil compared to the controls, and the cause was considered to be at least partly related to decreased water consumption. Hematological data showed significant increase of MCV in 800 ppm glycodil females and decrease of WBC in 3600 ppm oleate ester females. In serum biochemistry, increase of total cholesterol and potassium and decrease of ALT were detected in 800 ppm glycodil males, 3600 ppm linoleate ester males, and 800 ppm glycodil females, respectively. Serum creatinine levels in both sexes were decreased in the 800 ppm glycodil group. Relative weights of kidney and spleen were significantly increased in 200 and 800 ppm glycodil males and 800 ppm females. In addition, increase of relative kidney weights was also found in 3600 ppm oleate ester males. On histopathological assessment, increased cellularity of the renal cortex was observed in the epididymal ducts of 800 ppm males, but not in ester groups. Although more detailed analysis will be needed to clarify any testicular toxicity of glycodil and in vivo genotoxicity of GEs, our results suggest that oleate and linoleate esters might be less toxic to F344 rats than glycodil itself.

**2341 Modes of Action Underlying Citrinin-Induced Renal Carcinogenesis.**

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Citrinin (CTN), a mycotoxin produced by Penicillium and Aspergillus, is known to induce renal tumors in rats; however, the involvement of genotoxic mechanisms remains unclear. To evaluate the genotoxic potential of CTN, reporter gene mutation, comet, and micronucleus assays were performed. For the reporter gene mutation assay, groups of 5 male F344 rats were treated with CTN at the same doses by gavage for 2 days to extirpate the kidneys 3 days after the last dosing. For the comet and micronucleus assays, groups of 5 male F344 rats were treated with CTN at the same doses by gavage for 2 days to extirpate the kidneys or bone marrow, respectively. In the reporter gene mutation assay, the high dose (40 mg/kg) was decreased to 30 mg/kg on day 4 because of severe weight loss. The results of the reporter gene mutation and comet assays suggested that CTN did not induce DNA damage and subsequent gene mutations. Positive result was obtained only in the micronucleus assay, which might result from numerical chromosomal aberrations due to microtubule dysfunction by CTN. Therefore, it seems likely that non-genotoxic mechanisms are involved in CTN-induced carcinogenesis. In kidney samples from gpt delta rats, increases in the labeling indices of proliferating cell nuclear antigen (PCNA)-positive cells and mRNA expression levels of cell cycle-related genes (i.e., cyclin E1, cyclin A2, cyclin B1, and EZF1) were observed at all doses, despite the fact that the low dose showed no toxicological effects. Accordingly, the promotion of cell cycle progression observed in the kidneys of CTN-treated rats may have resulted from a direct mitogenic function of CTN. Increased phospho-ERK levels resulted from a direct mitogenic function of CTN. Increased phospho-ERK levels.

**2342 The Dynamics of a Harmful Algal Bloom and Paralytic Shellfish Toxins in Juneau, Alaska.**


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Paralytic shellfish poisoning (PSP) is a deadly neurological syndrome resulting from the ingestion of shellfish containing high levels of paralytic shellfish toxins (PSTs), and approximately seven cases of PSP are reported in Alaska annually. The main cause of PSTs is the neurotoxin saxitoxin, which is produced by the dinoflagellate *Alexandrium*. In June and July 2012 during this summer undergraduate project at the University of Alaska Southeast, plankton tows, seawater samples, and bivalve samples revealed a significant bloom of *Alexandrium* and subsequent toxin event in seawater and bivalves. Saxitoxin was extracted from the water column by filtering 1L of seawater and sonicating the filter for one hour in diH2O. The concentration of saxitoxin in seawater, measured using ELISA, ranged from 26 ug/L to 230 ug/L during the bloom. Nine species of bivalves, *Saxidomus giganteus*, *Mytilus trossulus*, *Clinocardium nutattles*, *Protocochlora staminea*, *Mya truncata*, *Mya arenaria*, *Tresus capax*, *Macromeris polyomma*, and *Hiatella arctica*, were sampled for saxitoxins using ELISA following soft tissue homogenization and extraction. Two species, *Mytilus trossulus* and *Saxidomus giganteus*, exceeded the regulatory limit for saxitoxin, 80 ug/100g wet wt . *Mytilus trossulus* saxitoxin concentrations ranged from 67 to 215 ug/100g, while *S. giganteus* ranged from 143 to 279 ug/100g. After the bloom, concentration of saxitoxin in the seawater and *M. trossulus* decreased, at a rate of 5.7ug/day, but remained elevated in *S. giganteus*. The concentration of saxitoxin in *M. trossulus* was below the FDA regulatory limit of 80ug/100g wet wt. 26 days after the bloom of *Alexandrium* was detected. Recreational harvest of bivalves in Alaska is not regulated for PSP; however, these results suggest that PSP risk is very high, particularly for *S. giganteus*.

**2343 Acute and 28-Day Oral Toxicity Evaluation of siRNAs and Longer Double-Stranded RNAs in Mice.**


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RNA interference is being used in agricultural biotechnology as a selective tool for developing crop traits. There are numerous biological barriers to uptake and activity of ingested nucleic acids that are ubiquitous components of animal diets. To evaluate the potential for adverse effects of dietary double stranded RNAs (dsRNAs), we conducted oral toxicity studies in mice with a pool of four 21-mer small interfering RNAs (siRNAs) and a 218 base pair dsRNA targeting the mouse ortholog of vacuolar ATPase (vATPase). When dsRNA targeting the insect ortholog of vATPase is expressed in corn plants, they are insecticidal against corn rootworm. Test materials were administered to CD-1 mice by oral gavage in a single dose acute toxicity study at 2000 mg/kg and in a 28 day repeat dose oral toxicity study at 1, 10, and 100 mg/kg. Toralai yeast RNA was included as a control in both studies. There was no impact of treatment on body weight, food consumption, clinical observations, or gross pathology in the acute toxicity study: The acute NOAELs for both the siRNAs and dsRNA were 2000 mg/kg, the highest doses tested. In the 28-day study, there were no treatment-related adverse effects on body weight, food consumption, clinical observations, clinical chemistry, hematology, gross pathology, or micropathology. The NOAELs in the 28 day study for both the siRNAs and dsRNA were 100 mg/kg, the highest doses tested. In summary, siRNAs and longer dsRNAs with 100% sequence identity to mouse vATPase do not result in adverse effects when administered orally to mice in a large dose. These results are consistent with the current body of knowledge that exogenous dsRNA molecules in food, even those with sequences identical to human or animal genomes, are safely consumed.
Arsenic is a ubiquitous, naturally occurring metalloid that poses a significant human cancer risk. While water consumption provides the majority of human exposure to arsenic, naturally occurring levels of arsenic in lakes, grains, vegetables, meats and fish, as well as through processed water containing arsenic (e.g. cooking rice) present a significant exposure to millions of individuals worldwide. To estimate the global burden of diseases attributable to toxic inorganic arsenic in food, we first evaluated the weight of evidence that supports a causal role for arsenic in a number of cancers and non-cancer disease endpoints. We determined that there was substantial evidence from large epidemiological studies that arsenic causes skin, lung, and bladder cancer in humans. The body burden of toxic arslenicals from foods is difficult to estimate and highly variable due to the natural distribution of arsenic in soils and water and the complication posed by multiple toxic inorganic and organic arslenicals, as well as non-toxic organic arslenicals contributing to total arsenic levels. Therefore we used GEMS/FAO-STAT estimates of food consumption in thirteen global clusters and JECFA reported measurements of total and inorganic arsenic in different foods to determine the upper and lower boundaries of foodborne inorganic arsenic exposures. We converted previously reported slope parameters from an Alternative Synthetic Route.

The present study evaluated the mutagenic and toxicologic potential of a proprietary analogues are metabolically activated in a similar fashion as the parent furan, yielding trans-Resveratrol is a naturally-occurring, polyphenolic compound found predominanly in grapes and red wine, and resVida (≥ 99.0% trans-resveratrol manufactured by DSM) has previously been assessed for safety based on toxicity studies (Williams et al, 2009) and obtained self-GRAS status in 2008. An alternative manufacturing process is now available in which trans-resveratrolyl is obtained with the same high purity of ≥ 99.0% but with different trace components to those in the original manufacturing process. Several of these new resveratrol-related by-products are found in nature (for example trans-pterostilbene). A safety assessment of the new process trans-resveratrol and its by-products was undertaken. The assessment process comprised:

- Identification of the new by-products
- Literature review for information on new by-products
- In silico analysis (DEREK) for toxic alerts of new by-products
- Structural analog comparison where appropriate.

- Ames tests (Salmonella Typhimurium Reverse Mutation Assays) with representative and spiked batches (to maximal specification level) of the new process material

- Definition of a new upper limit specification for newly identified by-products

Information presented from this process includes summarized data from the Ames test, which showed no mutagenic potential, with or without S9. From the assessment of all this information it was concluded that a ≥ 99.0% trans-resveratrol produced from the alternative route is safe and suitable for use within the marketing limits defined in the GRAS evaluation of the original process material.


In thermally treated products, a series of alkylated furan derivatives have been found, in particular 2-substituted alkylfuranfurans such as 2-methylfuran. These methyl analogues are metabolically activated in a similar fashion as the parent furan, yielding highly reactive unsaturated dialdehydes. There is limited toxicological data available for 2-methyl furan which makes conducting a risk assessment difficult. This pilot study was designed to determine the dermal uptake of 2-methylfuran for future subchronic studies needed to determine a NOAEL. Male Fischer 344 rats 5-6 weeks of age were administered 2-methylfuran by gavage to final concentrations of 0, 0.4, 1.5, 5, 6, 12, or 25 mg/kg bw/day. The animals were weighed daily prior to gavage and food consumption was measured weekly. The liver was the primary target organ which developed dose-dependent toxicity. Relative liver weights were increased by 42% at 25 mg/kg bw/day. Histological changes in the liver were observed at 0.4, 1.5, 3, 6, 12 and 25 mg/kg bw/day. These changes were not accompanied by clinical changes in serum liver enzyme markers such as ALT, ALP and AST. Clinical biochemistry markers for kidney were altered but these were not accompanied by histological changes. At 25 mg/kg bw/day, spleen weights were increased and the prostate was significantly increased in size. Some hematological parameters were also altered. In this pilot study, the liver was the major target organ.
for 2-methylfuran as indicated by changes in gross, histological and clinical parameters. Although the weights of the organs including prostate, kidneys and spleen, these were not accompanied by histological changes. The results of this study will be used to conduct a future subchronic study to establish a NOAEL for risk assessment purposes.

2349 Characterization of Bacterial Mutagenicity QSAR Predictions of Food Additives to Support Safety Assessments in a Regulatory Setting.
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Assessment of food additive safety at the U.S. FDA has utilized quantitative SAR (QSAR) analysis models for endpoints ranging from genetic toxicity, reproductive and developmental toxicity to rodent carcinogenicity. However, many of the QSAR models in routine use were originally developed for drug and industrial chemical assessments. Consequently, a performance assessment was undertaken on their use for compounds of interest to FDA CFSPAN and the performance of a QSAR model developed for predicting overall Salmonella mutagenicity. The test set was used to assess the suitability of the chemical space of the model for predicting food additive compounds. Overall performance statistics for accuracy, sensitivity, specificity and domain of applicability were measured. Structural classes were identified through compound clustering based on structural fingerprints that were well-predicted, poorly predicted, and not able to be predicted. Additionally, a set of compound structural alerts that represented different mutagenic toxicophores was assembled and used to help more precisely quantify performance. Variation in performance was observed across different toxicophores, providing a detailed picture of the model's strengths and weaknesses from a structural perspective in assessing food additives.

2350 The Role of Palmitoylation in Chemical and Microbial Toxicity: Signal Pathways, Protein Binding and Trafficking.
Multicellular organisms use chemical messengers to transmit signals among organelles and to other cells. Relatively small hydrophobic molecules such as lipids are excellent candidates for this signaling purpose. In most proteins, palmitic acid and other saturated and some unsaturated fatty acids are esterified to the free thiol of cysteines and to the N-amide terminal. This process enhances the surface hydrophobicity and membrane affinity of protein substrates and play important roles in modulating protein trafficking, stability, and sorting. Protein palmitoylation has been involved in numerous cellular processes, including signaling, apoptosis, and neuronal transmission. The palmitoylation process is involved in diseases such as Huntington's disease, various cardiovascular and T-cell mediated immune disorders, and cancer. Our study on lipopolysaccharide and deoxyxynivalenol treatment to rats provides insights on the role of protein palmitoylation in chemical and microbial toxicity. In the liver of animals treated with 10 mg/kg DON, palmitic acid decreased by 22% between 3 and 24 hr and increased 24% between 24 and 72 hr as compared to the controls. LPS at 85 μg/kg caused 54% decrease in elatic acid between 3 and 24 hr, and 7% between 24 hr and 72 hr while stearic acid decreased 33% between 3 and 24 hr and 60% between 24 and 72 hr as compared to the controls. Palmitate is a component of the LPS of Gram-negative bacteria. The bacterial outer membrane enzyme lipid A palmitoyltransferase PspG confers resistance to host immune defenses by transferring a palmitate chain from a phospholipid to the lipid A component of LPS. PspG is sensitive to cationic antimicrobial peptides (CAMP) which are included among the products of the Toll-like receptor 4 (TLR4) signal transduction pathway. This modification of lipid A with a palmitate appears to both protect the pathogenic bacteria from host immune defenses and attenuate the activation of those same defenses through the TLR4 signal transduction pathway.

2351 Compartment-Regulated Expression of Macrophage-Inhibitory Cytokine 1 under Murosal ER Stress.
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Endoplasm reticulum (ER) stress causes global translational arrest during protein biosynthesis. In spite of global translational arrest, the critical stress responsive genes, including macrophage inhibitory cytokine 1 (MIC-1), are particularly turned on. Functionally, MIC-1 played pivotal roles in ER stress-linked apoptotic death, which was also influenced by CEBP homologous protein, a well-known apoptotic mediator of ER stress. ER stress enhanced MIC-1 mRNA stability instead of transcriptional activation, and there were two mechanistic translocations critical for mRNA stabilization. First, C/EBP homologous protein triggered protein kinase C-linked cytosolic translocation of the HuR/EFLAV1 (Elav-like RNA-binding protein 1) RNA-binding protein, which bound to and stabilized MIC-1 transcript. As the second critical compartment-regulated modulation, ER stress activated ERK1/2 signals contributed to enhanced stabilization of MIC-1 transcript by controlling the extended holding of the nucleated mRNA in the stress granules fusing with the mRNA-decaying processing body. Taken together, these two sequential compartment-associated actions can account for stabilized transcription and subsequent re-initiation of translation of pro-apoptotic MIC-1 gene under mucosal ER stress (This study was carried out with the support of National Joint Agricultural Research Project of RDA (project number P008405932012) RDA, Republic of Korea).

2352 Azathioprine-Induced Hepatotoxicity in an In Vitro Inflammation-Immune Model.
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Azathioprine (AZP) is widely used in clinical practice for preventing graft rejection in organ transplantations, various autoimmune and dermatological diseases with documented unpredictable toxicity. Several experimental models suggested that an episode of inflammation during drug treatment predisposes animals to tissue injury. Inflammation caused by infections or endotoxins markedly activates NADPH oxidase. In the phagosome, superoxide radicals spontaneously form hydrogen peroxide (H2O2) and other reactive oxygen species. The effect of inflammation on AZP using “Accelerated Cytotoxicity Mechanism Screening” technique was investigated in this study. The concentration of AZP required to cause 50% cytotoxicity in 2 hr towards isolated rat hepatocytes was found to be 400 μM. AZP (400 μM) significantly increased cytotoxicity compared to control hepatocytes. When a non-toxic H2O2 generating system (glucose/glucose oxidase) was added to the hepatocytes prior to the addition of AZP, an increase in AZP cytotoxicity was observed. Because neutrophils or Kupffer cells release myeloperoxidase on activation, the effect of adding peroxidase to the hepatocytes exposed to H2O2 on AZP was also investigated. AZP showed a significant increase in cytotoxicity compared to drug-control in presence of glucose/glucose oxidase with or without horseradish peroxidase. A significant increase was also observed with glutathione depleted and catalase inhibited hepatocytes. Furthermore, AZP increased reactive oxygen species (ROS) formation, lipid peroxidation and decreased %mitochondrial membrane potential with our inflammation-immune model indicating the involvement of oxidative stress by glutathione oxidation, lipid peroxidation and mitochondrial toxicity. Protection was achieved by a ROS scavenger, 4-hydroxy-2,2′,6,6′-tetramethylpiperidene-1-oxyl (200 μM) and an antioxidant N,N′-diphenyl-p-phenylenediamine (2 μM). These results raise the possibility that the presence or absence of inflammation may be another susceptibility factor for azathioprine-induced hepatotoxicity.

2353 Indole-3-Carbinol and 3, 3′-Diindolylmethane Decrease Histone Deacetylase 3 Which Plays An Important Role in the Promotion of Staphylococcal Entertoxin B-Induced Immune Cell Activation.
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Staphylococcal enterotoxin B (SEB) is an exotoxin produced by the Staphylococcus aureus bacterium. This toxin is classified as a “superantigen” because of its ability to directly bind T cell receptors with MHC II class receptors of antigen presenting cells, which activates a large proportion of T cells. SEB is commonly associated with classic food poisoning, and more recently gained attention as a potential biological warfare agent since it is easily aerosolized. We have shown that indole-3-carbinol (I3C) and one of its byproducts, 3,3′-diindolylmethane (DIM), which are found in cruciferous vegetables, is able to reduce the number of SEB-activated T cells both in vitro and in vivo. These compounds were also able to reduce immune cell activation, induce apoptosis, and decrease proinflammatory cytokine release. In the current study, we assessed the role histone deacetylases (HDACs) played in SEB-treated cells, as well as what effect I3C/DIM had on them. Using antibodies specific for Class I or Class II HDACs, we showed that inhibition of Class II HDACs leads to decreased immune cell activation, increased apoptosis, and reduction in proinflammatory cytokine release in SEB-activated cells. However, inhibition of Class II
HDACs had opposite effects, suggesting a dual role of these HDAC classes in SEB stimulation, where Class I HDACs were important in SEB-mediated immune cell activation. Screening HDAC expression with western blots, we were able to determine that HDAC3 was the main HDAC upregulated after SEB stimulation, and I3C and DIM treatment was able to decrease this expression level. This research establishes for the first time the important role Class I HDACs, particularly HDAC3, play in SEB-induced immune cell activation. We were also able to provide more evidence for the effectiveness of I3C/DIM treatment in SEB through the downregulation of HDAC3. (Supported in part by NIH grants P01AT003961, R01AT006888, R01ES019313, R01MH094775, P20RR026368 and VA Merit Award BX001357).

Cytokines are important inflammatory mediators. Disturbance of the balance of pro- and anti-inflammatory cytokines may result in multiple organ toxicity. In this study, we compared two commonly used immunosassays for the detection of cytokines, i.e. Enzyme-Linked ImmunoSorbent Assay (ELISA) and Cytometric Bead Array (CBA). BD Oprelia ELISA kits and BD CBA Flex Sets were used to determine interleukin 4 (IL-4), interleukin 10 (IL-10) and interferon gamma (IFN-γ) levels in serum of untreated rats and in assay diluent. The intra-assay variation was determined after spiking with the individual recombinant standard of an ELISA kit at concentrations within the ELISA standard range or spiking with the combined recombinant standards of the CBA sets at concentrations within the CBA standard range. In addition, inter-assay variation was determined by spiking at two concentrations within the range of both ELISA and CBA standards and by analyzing these spiked samples using both methods on the same day. Both ELISA and CBA methods showed similar coefficients of variation for all cytokines, i.e. less than 15% for the majority of the measurements. The accuracy of assay diluent spiked with one or more cytokines was within 75-125% for all measurements using both methods. Spiking of serum samples showed clearly that serum contains factors that interfered with the quantification of IL-4, IL-10 and IFN-γ using the selected ELISA kits and CBA sets. For example, quantification of IL-10 in spiked serum showed an accuracy of approximately 30% using both methods and IFN-γ analysis resulted in an accuracy of <30% using ELISA and an accuracy of 70-80% using CBA. The absolute values of cytokines in spiked serum samples may differ between both methods, like for IFN-γ, but relative levels of cytokines always correlated with the spiked concentrations using either one of these methods. In conclusion, both ELISA and CBA showed similar precision and consistency in relative cytokine levels.

Rationale: The aryl hydrocarbon receptor (AhR) has emerged as an endogenous suppressor of cyclooxygenase-2 (Cox-2), Cox-2 is an immediate-early gene that is robustly increased by cigarette smoke exposure. We have published that the AhR suppresses cigarette smoke-induced Cox-2 protein but not mRNA, suggesting post-transcriptional regulation as a mechanism. The AhR may destabilize Cox-2 mRNA by retaining the RNA-binding protein (RBP) HuR in the nucleus. There is no known association between the AhR and HuR. Therefore, we investigated whether AhR-dependent retention of nuclear HuR is responsible for Cox-2 mRNA destabilization.

Methods: AhR+/−, AhR+/+, AhRDBD/DBD (harboring a mutant AhR unable to bind DNA) and AhRDBD/B6 mouse lung fibroblasts were exposed to cigarette smoke extract (CSE) for 3 h followed by Actinomycin D (ActD) for 30 minutes, 1 or 3 h, Cox-2 protein and mRNA were analyzed by western blot and qRT-PCR, respectively. HuR expression was assessed by western blot and immunofluorescence. AhR+/− cells were transfected with HuR siRNA and exposed to 1% CSE for 3 h with or without ActD for an additional 3 h. Cox-2 mRNA was then assessed by qRT-PCR.

Results: Steady-state Cox-2 mRNA levels significantly declined upon ActD treatment in AhR+/− cells, AhRDBD/DBD and AhRDBD/B6 cells, suggesting that the AhR destabilizes Cox-2 mRNA by a DRE-independent mechanism. Cox-2 mRNA instability was due to the nuclear retention of HuR. CSE did not alter HuR expression, but induced cytokines that HuR shuttles only in AhR+/− cells. Knockdown HuR in AhR+/− cells significantly decreased Cox-2 mRNA expression after exposure to ActD.

Conclusions: AhR-dependent retention of nuclear HuR suppresses cigarette smoke-induced Cox-2 protein by a mechanism that is independent of DNA-binding activity. These important findings open the possibility that a DRE-independent AhR pathway may be exploited therapeutically as an anti-inflammatory target.

Ultradextrous radiation (UVB) is the leading cause of skin cancer worldwide. UVB also modulates certain inflammation driven cutaneous pathologies such as contact hypersensitivity through actions on skin resident dendritic cell (DC) subsets. Transforming Growth Factor-β1 (TGF-β1) is a potent immunoregulatory cytokine in the skin microenvironment. Here, we show that TGF-β1 is required for UVB-induced activation and migration of dendritic cells to the skin draining lymph nodes. We indicated skin of Skin-Flax (SKFL) mice with UVB in the absence or absence of SB431542, a small molecule inhibitor of the TGF-β type I receptor and measured lymph node migration of skin dendritic cell subsets at acute time points. Topical inhibition of TGF-β1 pathway with SB431542 suppressed the migration of skin dendritic cell subsets, primarily CD103+ CD207+ and CD207- DC populations to the lymph nodes in response to UVB irradiation. In addition, in an ex vivo, skin explant assay for the migration of dendritic cells, UVB-induced DC migration into culture media was suppressed with topical inhibition with SB431542. In mice expressing a dominant negative receptor for TGF-β in CD11c+ dendritic cells (CD11c/TGF-βR1 DNR), UVB-induced migration of the DC subsets was suppressed directly linking TGF-β signaling in DC to UVB induced migration of DCs. Treatment with SB431542 also suppressed UVB-induced Interferon γ (IFNγ) secretion as well as the effector differentiation of T lymphocytes within the lymph nodes. Consistent with decreased activation within the lymph nodes, SB431542 decreased UVB activation of the skin infiltrating CD4 and CD8 lymphocytes after acute treatments and in UVB-induced skin tumors. Together, these data show that the TGF-β1 signaling pathway is important for the initiation of the inflammatory response to UVB irradiation of the skin, mediated primarily through the dendritic cells.

2356 Ultraviolet Radiation (UVB)-Induced Migration of Skin Dendritic Cell Subsets Is Mediated through Transforming Growth Factor Beta Signaling.

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Infection and inflammatory signaling can significantly alter drug metabolism enzyme expression (DME), thereby impacting the capacity of the liver to clear toxicants from the body. Previous work in our laboratory has detailed the modulation of gene expression of several hepatic DMEs during colonic infection with live Citrobacter rodentium (C. rodentium). These alterations included particularly strong downregulation of Fmo3 and Cyp4a family members, and appear to be largely independent of LPS-TRIF signaling. In order to elucidate potential signaling networks and pathways involved in this downregulation, we examined the impact of C. rodentium infection on the hepatic transcriptome using the Illumina MouseRef-8 v2 expression BeadChip. HeJ mice (n=4) lacking toll-like receptor 4 (TLR4), along with appropriate wild type animals (HeOUJ), were orally inoculated with live C. rodentium in a sucrose solution or received sterile sucrose. After 7 days, animals were sacrificed, livers were harvested, and RNA was prepared and submitted for analysis. Genes showing differential expression during infection were identified and pathway analysis using gene-set enrichment was performed. Increased numbers of genes with altered expression were found in HeJ mice as compared to HeOUJ controls. Several DME genes not previously reported as being altered in C. rodentium infection on the hepatic transcriptome using the Illumina MouseRef-8 v2 expression BeadChip.

2357 Alterations in the Hepatic Transcriptome during Live Citrobacter Rodentium Infection.

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Infection and inflammatory signaling can significantly alter drug metabolism enzyme expression (DME), thereby impacting the capacity of the liver to clear toxicants from the body. Previous work in our laboratory has detailed the modulation of gene expression of several hepatic DMEs during colonic infection with live Citrobacter rodentium (C. rodentium). These alterations included particularly strong downregulation of Fmo3 and Cyp4a family members, and appear to be largely independent of LPS-TRIF signaling. In order to elucidate potential signaling networks and pathways involved in this downregulation, we examined the impact of C. rodentium infection on the hepatic transcriptome using the Illumina MouseRef-8 v2 expression BeadChip. HeJ mice (n=4) lacking toll-like receptor 4 (TLR4), along with appropriate wild type animals (HeOUJ), were orally inoculated with live C. rodentium in a sucrose solution or received sterile sucrose. After 7 days, animals were sacrificed, livers were harvested, and RNA was prepared and submitted for analysis. Genes showing differential expression during infection were identified and pathway analysis using gene-set enrichment was performed. Increased numbers of genes with altered expression were found in HeJ mice as compared to HeOUJ controls. Several DME genes not previously reported as being altered in C. rodentium infection were identified, comprising P450s, UGTs and GSTs. Ontology term/pathways with high enrichment included Drug Metabolism, Mitochondrion, Oxidation Reduction, and Inflammatory Response. Other terms/pathways of interest include Apoptosis, Glutathione Transferase Activity, and Lipid Metabolism. An analysis of the potential transcription control pathways upstream of these observed gene expression changes was also performed. Potential upstream factors include AhR, ARNT, Ekl-1, and Hnf-3beta, among others. Taken together, these results demonstrate that drug metabolism gene expression is particularly sensitive to infection, and indicate several potential pathways that may be responsible for these effects. Supported by grant R01072372 from the NIH.
Silencing of Keap1 in Macrophages Boosts Lipopolysaccharide-Induced Transcription of Interleukin 6 via IkBα Activation.


Interleukin-6 (IL6) is a multifunctional cytokine that regulates immune and inflammatory responses. Multiple transcription factors, including NF-κB and nuclear factor E2-related factor 2 (Nrf2), are implicated in the transcriptional regulation of IL6. Kelch-like ECH-associated protein 1 (Keap1) is a substrate adaptor protein for a Cullin 3-dependent E3 ubiquitin ligase complex, which regulates the degradation of various vital proteins, including Nrf2 and IkBα. In agreement with previous studies, stable knockdown of Nrf2 in RAW 264.7 mouse macrophages led to significantly attenuated antioxidant response and decreased expression of IL6 under basal and lipopolysaccharides (LPS)-treated conditions. However, Nrf2 activation alone (e.g., under tert-butylihydroquinone exposure) did not increase the expression of IL6, suggesting that Nrf2 is a necessary, but not sufficient, factor in regulating LPS-induced transactivation of IL6. In contrast, silencing of Keap1 in RAW cells and human monocyte THP1 cells markedly augmented the expression of IL6 under non-stressed and LPS-challenged conditions. The enhanced expression of IL6 in Keap1-knockdown (Keap1-KD) cells was significantly attenuated by silencing of Ikβα, but not Nrf2, suggesting that stabilized Ikβα resulting from Keap1 silencing is the major downstream event responsible for the transactivation of IL6. This finding was further confirmed by the enhanced protein levels of IkBα and subsequent increased expression and phosphorylation of NF-κB p65 in the Keap1-KD cells. Together, the present studies demonstrated that silencing of Keap1 in macrophages boosts LPS-induced transcription of IL6 via IkBα activation. Given the importance of IL6 in inflammatory response, targeting Keap1 could be a novel approach in the treatment and prevention of inflammation and associated disorders.

Potent Protection against PM2.5 Diesel Exhaust Particle-Induced ROS Generation and Vasculature Permeability through Regulation of Nrf2-Induced Pathways by Triterpenoids.

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Epidemiologies suggest that an increase of PM2.5 diesel exhaust particles (DEP) in ambient air corresponds to an increase in myocardial infarctions within 48 hours. To cause such disorder, the close association of capillaries and alveoli should allow DEP traveling in the bloodstream, two inhaled DEP to get in close proximity to capillary endothelial tubes. However, the production of oxidative stress is considered to be one of the key mechanisms how particles affect tissues. Complex in vitro coculture systems may be valuable tools to study related processes and to further evaluate the effects of particles on the lung. Exposure to fine and ultra-fine ambient particles is still a problem of concern in many industrialised parts of the world and the intensified use of nanotechnology may further increase exposure to small particles. Among the various mechanisms, the production of oxidative stress is considered to be one of the key mechanisms how particles affect tissues. Complex in vitro coculture systems may be valuable tools to study related processes and to further evaluate the effects of particles on the lung (Klein et al., 2011). Therefore, a system consisting of four different human lung lines that should mimic the cell response of the alveolar surface in vitro was developed in order to be used with a native aerosol exposure system (Vitrocell™ chamber). It is composed of one alveolar type-II cell line (A549), differentiated macrophage-like cells (THP-1), mast cells (HMC-1) and endothelial cells (EA.hy 926), seeded in a 3D orientation on microporous membranes. Exposure to diesel exhaust could induce the cells with 2,2'-azobisis-2-methylpropanimidamide, dihydrochloride (AAPH; 20 mM), and quantified as the oxidative stress. Results are reported as fold increase in ROS production relatively compared to untreated cells. Single cell cultures of EA.hy 926 (11.8 +/- 1.4), THP-1 (11.5 +/- 3.3) and HMC-1 (14.7 +/- 2.9) showed significantly higher oxidative stress than the tetraculture (6.6 +/- 0.75). A549 cells showed the highest amount of oxidative stress (3.4 +/- 0.18) compared to other cultures. The interplay of model cell for the immune system (THP-1 and HMC-1) with A549 epithelial cells strongly influences the behaviour of our system, resulting in an alleviative effect for oxidative stress compared to the monocytes. The use of the tetraculture system may lead to a more realistic judgement about the hazard of new compounds in the future.

Differential Responses upon Inhalation Exposure to Biodiesel versus Diesel Exhaust on Oxidative Stress, Inflammatory, and Immune Outcomes.

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Biodiesel (BD) exhaust may have reduced adverse health effects due to lower mass emissions and reduced production of hazardous compounds compared to diesel exhaust. To investigate this possibility, we compared adverse effects in lungs and liver of BALB/c mice after inhalation exposure (0, 50, 150 and 500 μg/m3; 4 hr/day, 5 d/wk, for 4 wk) to combustion exhaust from 100% biodiesel (B100) and diesel (D100). Compared to D100, B100 exhaust caused a significant accumulation of oxidatively modified proteins (carbonyls), increase in 4-hydroxynonenal (4-HNE), reduction of protein thiol, depletion of antioxidant - glutathione (GSH), a dose-dependent increase in the levels of biomarkers of tissue damage (LDH) in lungs, and inflammation (myeloperoxidase, MPO) in both lungs and liver. B100 exposure also significantly enhanced expression of cytokines IL-6, and IL-12p70 (in a dose-dependent manner), along with IL-10, TNF-α and MCP-1 (increased compared to control) in both lung and liver tissues. Overall, the cytokine profiles in the lung and liver suggest that B100 and D100 exhaust elicit similar innate immune responses, predominantly involving T-cell independent pathways; however, the magnitude of inflammation was greater following B100 exhaust exposure. Interestingly, exposure to D100, but not B100 exhaust, induced a significant increase in the levels of IFN-γ in the lungs, suggesting a broader engagement of Th1 component by D100 exhaust. Based on this, we hypothesize that the distinctive organic compounds and/or oxidative products formed as a result of increased oxidative stress upon B100 exposure, are capable of targeting biological/molecular pathways that are distinct from D100 exposure. (This abstract does not represent US EPA policy).
Intranasal treatment with ferric oxide nanoparticles (α-Fe2O3 and γ-Fe2O3 NPs), in rats caused microglial proliferation and activation in olfactory bulbs, hippocampus and striatum. Our in vitro studies with SHSY5Y neuroblastoma cells exposed to 10 and 30 nm ferric oxide NPs showed over expression of alpha-synuclein protein, depletion of dopamine, and conditions for oxidative stress. Here, we examined the response of brain and liver free fatty acids (FFAs) in adult male Sprague-Dawley rats treated intraperitoneally (i.p.) either with saline (control) or ferric oxide (Fe2O3) – NPs at 25, 50 and 100 mg/kg. Rats were sacrificed 72 hrs after injection to harvest caudate nucleus and liver. Long chain FFAs were extracted with chloroform and methanol (4, 8 v/v) from tissue homogenates and the extracts were shaken, followed by centrifugation. The supernatants were reconstituted with Hepes, chloroform and methanol (3, 2, 4, 8 v/v). The chloroform was then evaporated under nitrogen. The residue was reconstituted with ether-hexane (50:50,v/v) and eluted by isocratic chromatography on a silica gel plate. FFAs were derivatized with BF3/methanol and fatty acid methyl esters were quantitated using gas chromatography. Concentrations of saturated FFAs (palmitic, stearic) in the liver and brain and did not change following injection of the iron NPs. However, unsaturated brain FFAs (oleic, linoleic) were decreasing in a dose-related fashion in the CN (p<0.05). The liver, the concentration of the unsaturated FFA's increased significantly at 25 and 50 mg/kg (p<0.05) but was no different from control at 100 mg/kg. These data indicate a differential response of liver and brain unsaturated fatty acids to iron nanoparticle exposure, suggesting different mechanisms in the liver and brain in response to oxidative stress.
toxicity of nanoparticles embedded in paints compared to pristine nanoparticles.


Nanomaterials are increasingly being used in the paint industry due to their unique physical and chemical properties. Nanoparticles often used in paints and coatings are TiO₂ (anti-UV, self-cleaning, air purification), Ag (anti-microbial) and SiO₂ (fire retardant, anti-scratch).

In this study, the toxic effects of 3 pristine nanoparticles (TiO₂, Ag and SiO₂), 3 aged paints containing nanoparticles (TiO₂, Ag and SiO₂) and control paints without nanoparticles were compared.

BALB/c mice were weekly oropharyngeally aspirated with nanoparticles or paint for 5 weeks. Mice were sacrificed 2 or 28 days after the last aspiration. The local (lung/bronchoalveolar lavage fluid) and systemic (blood) toxicity was evaluated (cell counts, inflammatory cytokines, blood clotting parameters).

The pristine nanoparticles showed no effects in the blood and a subtle toxic effect in the lungs, which was most pronounced in the case of Ag nanoparticles (increase in neutrophils (7.8±10³), 2-fold increase in pro-inflammatory cytokines KC and IL-1). The paints containing nanoparticles did not show significant toxicity.

In conclusion, we demonstrated that although pristine particles show some toxic effects, no significant toxicological changes were observed when they were embedded in a complex paint matrix.

A 15-Day Oral Exposure to Dispersed TiO₂ P25 Particles Induces Epithelial Barrier Dysfunction and Bacterial Translocation in the Rat Intestine.

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Aim: Titanium dioxide (TiO₂) has a long-standing use as food additive and is promised to be used in food packaging as antimicrobial in biosurfaced foods. Possible hazards of ingested TiO₂ particles for human digestive tract are under discussion. We addressed consequences for gut barrier function in rats orally exposed to TiO₂ P25 (85% anatase/15% rutile) at human level exposure. Methods: Male rats were orally given either vehicle (Ve) or TiO₂ P25 (provided by European Commission-Joint Research Center in the OECD sponsorship program) at 100, 2.5 mg/body for 28 days. After the last administration, we observed the localization and absorption of nSP via oral route in vivo and in vitro. BALB/c mice were weekly oropharyngeally aspirated with nanoparticles or paint for 5 weeks. Mice were sacrificed 2 or 28 days after the last aspiration. The local (lung/bronchoalveolar lavage fluid) and systemic (blood) toxicity was evaluated (cell counts, inflammatory cytokines, blood clotting parameters). The pristine nanoparticles showed no effects in the blood and a subtle toxic effect in the lungs, which was most pronounced in the case of Ag nanoparticles (increase in neutrophils (7.8±10³), 2-fold increase in pro-inflammatory cytokines KC and IL-1). The paints containing nanoparticles did not show significant toxicity.

In conclusion, we demonstrated that although pristine particles show some toxic effects, no significant toxicological changes were observed when they were embedded in a complex paint matrix.

The Effect of Nanoparticles from Secondhand Cigarette Smoke on the Mouse Lung.

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Second hand cigarette smoke (also named Environmental tobacco smoke (ETS)) is an environmental trigger factor that leads to airway inflammation and airway hyperresponsiveness (AHR) in susceptible individuals and animals. The constituents of ETS exist in the gas-phase and the aerosol particles which consist predominantly of nanoparticles (two dimensions less than 100 nanometres). The purpose of this study is to characterize the role of nanoparticles on ETS-induced airway responses.

The mice were exposed to side-steam tobacco smoke (SS), a surrogate to ETS, or 50 nm nanoparticles, or 80 nm nanoparticles, or gas-phase or filtered air (FA) for 3hrs. Lung function and inflammation in bronchoalveolar lavage (BAL) were measured following exposure. Methacholine (MCh) dose response for lung resistance (RL) was significantly elevated, and dynamic pulmonary compliance (Cdyn) was significantly decreased, in the SS, nanoparticles exposure groups compared with the FA and groups gas-phase exposure. At the same time, the total cells and neutrophils were significantly elevated in both SS and nanoparticles exposed mice. However, MCh dose response curves for RL and Cdyn, inflammation were not significantly changed in the 50 nm nanoparticles and 80 nm nanoparticles exposure groups.

These results suggest that nanoparticles from second hand cigarette smoke play an important role in smoking-induced lung injury.

The Comparative Immunotoxicity of Mesoporous Silica Nanoparticles and Colloidal Silica Nanoparticles in Mice.

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Mesoporous silica (MPS) nanoparticles (NP), which have unique pore structure, extremely high surface area and pore volume, have attracted attention for their potential biomedical applications, such as carriers for controlled drug delivery and matrix for tissue regeneration. To use MPS NPs for biomedical devices, their bio-compatibility both in vitro and in vivo should be confirmed. The lung surface area of NPs is one of the important determinants of toxicity such as cellular uptake and immune response. We previously first reported that MPS NPs exhibited less cytotoxicity and inflammation potential than general amorphous colloidal silica (Col) NPs on macrophages. However, the low cytotoxicity does not guarantee high biocompatibility in vivo. In this study, we compared in vivo immunotoxicity of MPS and Col NPs in mouse model to define the influence of pore structural conditions of silica NPs. Both MPS and Col NPs (2, 20, 50 mg/kg/day) were intraperitoneally administered in female BALB/c mice for 4 weeks. There was no overt sign of clinical toxicity in both MPS and Col treated mice. Interestingly, the in vivo test showed opposite results from in vitro. MPS NPs significantly increased weight of liver and spleen, and proliferation of splenocytes. MPS NPs treated mice showed the altered lymphocyte population (CD3+, CD45+, CD4+ and CD8+) of spleen, increased serum IgG and IgM levels, and histological changes. In spite of the slight changes in lymphocytes population of spleen, Col NPs did not alter other immunological factors. Our results showed that in vivo exposure of MPS NPs causes more damages in systemic immunity than Col NPs by the immunoenhancement of spleen. The in vivo data showed opposite results from in vitro showing less cytotoxicity of MPS NPs. Our results suggest the importance of confirmation of biocompatibility both in vitro and in vivo during the design of new nanomaterials. These findings may provide useful information for the bioapplication of silica NPs.

Evaluation of Intestinal Absorption of Amorphous Silica Nanoparticles.

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With the recent development of nanotechnology, amorphous silica nanoparticles (nSP) with particle size below 100 nm have already been used in various foods as antac ing agents. Therefore, to ensure the safety of nSP, it is an urgent need to obtain safety information of nSP. However, there is little information about biodistribution of nSP after oral administration and absorption. In this study, we examined biodistribution and absorption of nSP via oral route in vivo and in vitro. BALB/c mice were orally exposed to nSP with diameter of 70 nm (nSP70) or 1000 nm (nSP1000) at 2.5 mg/body for 28 days. After the last administration, we observed the localization of silica particles in tissues by transmission electron microscope. Both silica particles were observed in some tissues such as spleen and liver, although these results were qualitative analysis. Next, we evaluated the absorption of silica particles through intestine quantitatively by everted sac method. Although about 0.3% of nSP1000 in mucosal side was absorbed into serosal side, the level of absorbed nSP70 was about...
Conclusion: Antitumor activity of IndOH in brain is kept when it is combined to IndOEt in NC. Fluorescence in the parenchyma was also observed 20 minutes, 1 and 2 hours (P<0.05). The BBB permeability was measured by the Evans blue extravasation assay. Effect of two week-IndOH/IndOEt-NC treatment on the volume of a brain tumor (induced by direct injection of glioblastoma cell line [GL261] into the brain) was measured. Ethical Committee number: CEUA/FCF/349. Results: The intensity of fluorescence outside the vessels dramatically increased 30 minutes following i.v. injection compared to 10 minutes time (P<0.05). The interaction of leukocytes-rhodamine-labeled and platelets-FITC-labeled with the endothelial cells of pial venules. The BBB permeability was measured by the Evans blue extravasation assay. Effect of two week-IndOH/IndOEt-NC treatment on the volume of a brain tumor (induced by direct injection of glioblastoma cell line [GL261] into the brain) was measured. Ethical Committee number: CEUA/FCF/349. Results: The intensity of fluorescence outside the vessels dramatically increased 30 minutes following i.v. injection compared to 10 minutes time (P<0.05). No increase in leukocyte or platelet adhesion to endothelial cells, and Evans blue extravasation were noticed. IndOH/IndOEt-NC profoundly reduced the brain tumor. Conclusion: Antitumor activity of IndOH in brain is kept when it is combined to its ester into NC. Further studies are needed to study the safety and efficacy of this combination.

Supported by: FAPESP, CNPq.

Oxidative stress was indicated as one of the main mechanisms associated with nanoparticle (NP)-induced toxicity. In this study, the validity of plasma F2-isoprostanes (F2-isoPs), a proposed oxidative stress marker (Halliwell & Lee, Antioxid Redox Signal, 2010), was examined in rats treated intratracheally with a single dose of cadmium-doped silica nanoparticle (SiNP-Cd), a model nanoparticle previously shown to induce oxidative stress in vivo (Cocci et al, J Nanopart Res, 2012). The response to SiNP-Cd (1 mg/rat) was evaluated 2 hr, 7 and 30 days post-instillation by immunocytochemistry analysis of superoxide dismutase (SOD1), inducible nitric oxide synthase (iNOS) and cyclooxygenase type 2 (COX-2) expression in pulmonary tissue. Lung and plasma levels of F2-isoPs were measured in parallel by GC/NICI-MS/MS analysis. Furthermore, the effects of SiNP-Cd were evaluated in comparison with those caused by equivalent amounts of CdCl2 or SiNP. In the animals exposed to SiNP-Cd, pulmonary SOD1, iNOS, and COX-2 immunoreactivity was enhanced in a time-dependent manner (7 <30 days). Pulmonary total F2-isoPs were also increased significantly on thirty days post-exposure (46.7±11 ng/g in SiNP-Cd vs 32.8±7.8 ng/g in control). Pronounced elevation of free F2-isoPs similarly occurred in plasma (54.6±22 pg/ml in the SiNP-Cd group compared to 28±8 pg/ml in controls). The increase in plasma F2-isoPs was already detectable at day 7 and lasted until day 30 post-exposure. In the animals treated with silica nanoparticles no changes were observed regarding the immunohistochemical and biochemical parameters tested. The pulmonary response to CdCl2 was less pronounced than that found with SiNP-Cd. These results indicate (i) the potential of SiNP-Cd to cause long-lasting oxidative tissue injury following pulmonary exposure in rat and (ii) a promising role for F2-isoPs as indicator of nanoparticle-induced oxidative insult (Grants: Italian Ministry of Health & University, and CARIPLO Foundation Rf. 2011 – 2096).

The potential impact of nanoparticles on the environment and on human health has attracted considerable attention in recent years. Transcriptomics data generated from tissues or cells exposed to nanoparticles are being gathered at ever-increasing rates. In addition to the importance of the original findings, such data can have value if interpreted in a broader context and combined with other published results. To encourage the efficient use of the data, we have developed NanoMiner (http://nanominer.cs.tut.fi/), an integrative transcriptomics data resource for nanoparticle research. The data in NanoMiner is collected from public repositories, and the database currently contains 404 human transcriptomics samples of cells exposed to various types of nanoparticles. All samples in NanoMiner have been annotated, preprocessed and normalized using standard methods to ensure the quality of the data analyses and to enable systematic use of the database across different experimental setups and platforms. With NanoMiner, it is possible to: 1) search and plot the expression profiles of one or several genes of interest, 2) cluster the samples within the datasets, 3) find differentially expressed genes in various nanoparticle studies, 4) detect the nanoparticles causing differential expression of selected genes, 5) analyze enriched Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways and Gene Ontology (GO) terms for the detected genes, and 6) search the expression values and differential expressions of the genes belonging to a specific KEGG pathway or Gene Ontology. The NanoMiner database is thus a valuable collection of microarray data and can also be used as a data repository for future analyses.

Nanoscaled europium oxide (EuO3) particles were selected to investigate the bioenergetics following inhalation. The rare earth Eu allowed a very high accuracy in analysis of potential translocation from lungs to remote organs. An aqueous dispersion of commercially available EuO3 particles (0.1 w-%) was prepared in phosphate buffer (0.15 w-%) incl. bovine serum albumin (0.25 w-%). A suspension partially consisting of nanoscaled particles could be realized by mechanical homogenization and ultrasonic treatment and was aerosolized with pressurized air. Rats inhaled the dry aerosol for 6 hours in a single inhalation. Phosphate facilitated the disintegration of the EuO3 particles in lung alveoli after deposition. The potential translocation of EuO3 particles was followed by chemical Eu analysis and transmission electron microscopy (TEM). Using chemical analysis, 36.8 μg/lung EuO3 were detected 1 hour after inhalation in lungs. The amount declined slightly to 34.5 μg after 1 day and 35.0 μg after 5 days. The liver showed an increase of EuO3 from 32.3 ng 1 hour up to 294 ng 5 days after inhalation. Additionally, lung-associated lymph nodes, thymus, kidneys, heart, and testes exhibited an increase of EuO3 over the time period investigated. In the blood, the highest amount of EuO3 was found after 1 hour whereas feces, urine and mesenteric lymph nodes revealed the highest amount after 1 day. In the other organs such as brain, spleen, adrenals and epididymides no changes of the Eu amount were detected. By TEM analysis, EuO3 particles could be detected only in lungs, in liver, however, with one of the highest chemical Eu concentrations, no particles were detectable. In conclusion, mixed type metal oxide/phosphate particles are a suitable tool for biokinetic investigations after inhalative uptake. The use of EuO3, combined with chemical and TEM analysis was a very suitable model to examine the translocation potential. Bioavailability was limited to soluble EuO3, a translocation of EuO3 particles was not evident.

The potential impact of nanoparticles on the environment and on human health has attracted considerable attention in recent years. Transcriptomics data generated from tissues or cells exposed to nanoparticles are being gathered at ever-increasing rates. In addition to the importance of the original findings, such data can have value if interpreted in a broader context and combined with other published results. To encourage the efficient use of the data, we have developed NanoMiner (http://nanominer.cs.tut.fi/), an integrative transcriptomics data resource for nanoparticle research. The data in NanoMiner is collected from public repositories, and the database currently contains 404 human transcriptomics samples of cells exposed to various types of nanoparticles. All samples in NanoMiner have been annotated, preprocessed and normalized using standard methods to ensure the quality of the data analyses and to enable systematic use of the database across different experimental setups and platforms. With NanoMiner, it is possible to: 1) search and plot the expression profiles of one or several genes of interest, 2) cluster the samples within the datasets, 3) find differentially expressed genes in various nanoparticle studies, 4) detect the nanoparticles causing differential expression of selected genes, 5) analyze enriched Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways and Gene Ontology (GO) terms for the detected genes, and 6) search the expression values and differential expressions of the genes belonging to a specific KEGG pathway or Gene Ontology. The NanoMiner database is thus a valuable collection of microarray data and can also be used as a data repository for future analyses.

Nanoscaled silica (SiO2) nanoparticles are widely used in many diverse industries, such as polishing agents in semiconductor fabrication, as potential biomedical drug delivery agents, and in fabrics to make them wrinkle free. Our goal was to evaluate the ef-
effects of nano-silica (SiO2, longest dimension 51.9 ± 16.2 nm, aspect ratio 1.1 ± 0.1) on small arterioles in elderly, Wistar rats. First, a suspension of SiO2 (in 1gm/ml albumin, saline) was directly applied to arterioles (micropipette, 137 pg total dose), using a hamster cheek pouch intravital microscopy model (isoflurane, N=6). Endothelial dysfunction (loss of dilation to acetylcholine, ACH 10-4M) was evident within minutes (dilation of 78±14% to constriction -5±9%, p<0.05), while dilation to adenosine was unaffected. Constrictor responses to phenylephrine were diminished by SiO2 exposure (from -61±4% to -12±2%); importantly, the baseline diameters were not altered by direct exposure to SiO2, and the vessels retained some responses. Next, mice were exposed to 20ug of SiO2 via aspiration, and 24 hours later the cremaster m. model was examined (isoflurane). Db/db controls (N=6) have mild endothelial dysfunction seen as a diminished dilation to ACH (10-4M, dilation of 43±14% to 10±6%); Exposure to SiO2 (40gm wt, N=4) induced a profound endothelial dysfunction seen as constriction to ACH (-13±2%); Dilation to adenosine was unaffected. In C57BL/6, exposure to SiO2 (25gm wt, N=2) did not significantly alter dilation to ACH or adenosine compared to controls (N=6). Further, in the aspiration model, constrictor responses to phenylephrine were not affected in either strain. Thus, direct exposure to 137 pg SiO2 induced endothelial dysfunction immediately in healthy hamsters. Aspiration of 20ug induced a profound endothelial dysfunction in diabetics but not in the genetic background controls. (NIH DK68401, HL5492)

2377 Copper Nanoparticles or Ionic Copper (II) Causes Neurotoxicity and Cardiotoxicity in Zebrafish Embryos.

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Copper oxide nanoparticles (Cu-NPs) are frequently used in medical devices, paints, fabrics or as antimicrobials. Their industrial applications may lead to the contamination of aquatic ecosystems. The toxicological and human health risks of NPs in the environment are hard to evaluate due to a lack of knowledge about the mechanisms by which NPs interact with biological systems. In this study, we investigated the toxicity of Cu-NPs and the ionic copper(II) form in wild-type (WT) zebrafish (Danio rerio, AB-strain) embryos and hh9-GFP transgenic zebrafish (Danio rerio, AB-strain) embryos by comparing bare Cu-NPs to the mass equivalent ionic form of copper(II) (CuCl2) at various concentrations (1.25- to 20 ug/ml). The toxicity was determined by phenotypic changes in the zebrafish embryos including survival, heart rate, motor neuron development and apoptotic forfeiture. Both Cu-NPs and CuCl2 were lethal to zebrafish embryos at 20 ug/ml (within 24-hrs) and 10 ug/ml (within 48-hrs), with CuCl2 being more toxic at equivalent mass concentrations. Similarly, the heart rate was significantly reduced following exposure to either Cu-NPs or CuCl2 in a concentration-time dependent manner. Additionally, the embryo permeability studies showed that exposure to either Cu-NPs or CuCl2 (5 ug/ml) for 24-hrs significantly increased the topological absorption of the fluorescent tracer 6-coumarin (6CM). Furthermore, embryos treated with either Cu-NPs or CuCl2 (2.5 ug/ml for 48-hrs) showed a significant reduction (nearly 2-fold) in spinal motor neurons. These results indicate that both CuCl2 and Cu-NPs can be acute, and lethal to zebrafish embryos causing significant neurotoxicity and cardiotoxicity at exposure levels that do not cause lethality.

2378 Nickel Oxide Nanoparticles Provoke Intrinsic Apoptotic Pathway in HepG2 Cells, Male Wistar Rats and Tomato Seeding Roots.

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Many features of programmed cell death in plants resemble with those observed in human and animals. Therefore, this study has been done with a rationale to provide first evidence on the molecular toxicity of nickel oxide nanoparticles (NiO-NPs, <50 nm) in human colon adenocarcinoma (HepG2) cells, male Wistar rats and tomato seeding roots. The cytotoxicity studies with NRU and MTT assays in HepG2 cells revealed 20.6 % and 18.4 % decline in the cell survival at the highest concentration of 100 μg/ml. Treated cells showed an increase in intracellular ROS generation and 30.5 % increase in flow cytometric sub-G1 apoptotic peak at 100 μg/ml NiO-NPs. Quantitative real-time PCR analysis of apoptotic pathway genes (P53, caspases 3 and 9) showed 2.0, 1.2 and 1.1-fold higher expression in HepG2 cells. Furthermore, western blot experiments also showed the greater activity of P53 and caspase 3 genes in cells exposed to 100 μg/ml NO-NPs. Oral exposure of male Wistar rats with 1, 2 and 4 mg/kg b.w of NiO-NPs for 48h and 14 days showed increased intracellular ROS generation. DNA damage, micronuclei formation and apoptosis in bone marrow cells. Tomato seeds exposed to NiO-NPs for 4 h, exhibited repression of root length, higher activities of antioxidant enzymes, and increased frequency of apoptotic and necrotic cells in comet assay. Flow cytometric analysis of 2 mg/ml treatment group revealed 122% higher ROS generation with alteration of mitochondrial membrane. Cell cycle data showed a shift of 65.5% cells towards apoptotic subG1 phase vis-à-vis control showed 16.5% cell in subG1. An increase in caspase-3 like protease activity validates the involvement of mitochondrial dependent intrinsic apoptotic pathway. Thus, this study has provided a new insight into the fundamental mechanism of NO-NPs induced toxicity and signifies its potential to induce cell death in animal and plant cells.

2379 Quantifying Quantum Dots in Frozen Tissue Sections Using Autometallography.

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Quantum dots (QDs) are engineered nanoparticles frequently composed of a CdSe core, ZnS shell, and an assortment of polymer coatings specific to their application. QDs are used in electronic systems because of their semiconductor properties and in biomedical research and medicine as imaging tools because of their unique fluorescence properties. Their widespread use and heavy metal core composition have raised concerns about their safety. An important consideration in evaluating QD toxicity is the accurate quantification of these nanoparticles within tissues. The current measurement methods favored include inductively coupled plasma mass spectrometry (ICP-MS) for the metal components of QDs, and QD fluorescence directly in tissue sections using microscopy. However, ICP-MS is expensive and cannot distinguish between metals present in QDs or free ions, and fluorescence microscopy is often difficult because of interfering tissue autofluorescence. We adapted a silver-enhanced autometallography technique for detecting QDs in frozen tissue sections. This technique is efficient and inexpensive, and quantification of both metal imaging and spectrometry correlates well with direct QD fluorescence measurements. The ability to efficiently measure QDs in tissues will provide important dose information that can be used for evaluating the adverse health effects of QD exposures.

2380 Influence of Primary Particle Size and Agglomeration State of Inhaled Nano-TiO2 on Rat's Pulmonary Response.

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The physico-chemical properties of nanoparticles (NP) and their agglomeration state influence their toxicokinetics, reinforcing the importance of the characterization of the exposure dose. The objective of this study was to evaluate the influence of initial particle size and agglomeration state of inhaled 20 mg/m3 TiO2 aerosols on rat’s pulmonary response. Groups of rats (n=6) were exposed for 6 hrs to aerosols composed of either 5, 10-30 or 50 nm TiO2. Two distinct agglomeration states were obtained. Aerosols were composed majorly of either large (L) (>100 nm) or small agglomerates (SA) (<100 nm). A control group was exposed to compressed air. Exposures were characterized using weight measurement for mass concentration, an electrical low pressure impactor (Dekati) for size distribution and electron microscopy for agglomerates observation. Pulmonary response was analyzed 16 hr after the end of exposure through bronchoalveolar lavage fluids and lung histology. Total cell count, number of macrophages and neutrophils were increased statistically (p < 0.05) compared to control for the 10-30 and 50 nm LA aerosols, while increases were significant only for total cell count and number of macrophages for the 5 and 10-30 nm SA aerosols. For each initial particle size, percentages of particulate laden macrophages in LA aerosols were higher and statistically different from the SA ones. Morphological assessments of lung tissue showed that cellular infiltrates were more important in exposed groups compared to controls, with exemption of the 50 nm SA group. Our results indicate that higher increases in neutrophil number are related to LA aerosols, as for initial particle size, 10-30 nm TiO2 NP seemed to induce more pronounced effects. Overall, this suggests that initial particle size and agglomeration state play a key role in the toxicity of TiO2 aerosols.
Silica nanoparticles (SiO2-NPs) are the one of most widely used and important nanomaterials in nanotechnology. Lung tissue is one of the main routes of entry nanoparticles, which may cause severe pulmonary toxicity. However, the toxicological effects and the precise mechanisms of SiO2-NPs on lung are still unclear. Here, we attempted to investigate the toxic injuries and the definitive mechanism of SiO2-NPs on the acute pulmonary toxicity. The adult male ICR mice were exposed to intratracheal dose of 50 mg/kg SiO2-NPs and lung tissue were collected after 7 days. Our results found that SiO2-NPs increased 40% mortality rate and significantly induced pulmonary morphological and histological changes with neutrophils, macrophage and fibroblast cells from the terminal bronchial. The lung tissue weight/body weight ratio (LW/BW) increased 2-fold suggested that SiO2-NPs may trigger pulmonary edema. Meanwhile, the malondialdehyde (MDA) levels in the treated lung tissue were increased. Moreover, SiO2-NPs caused apoptosis-related signals, including up-regulation of Bax and down-regulation of Bcl-2 and activation of caspase cascades mRNA expression, which accompanied with triggered the endoplasmic reticulum (ER) stress identified through several key molecules, such as activating the CHOP, XBP-1, caspase-12, and increasing the GRP-78/94 mRNA expression. These results suggest that SiO2-NPs induced an oxidative stress, and cause acute pulmonary toxicity through mitochondria and endoplasmic reticulum pathways.
2385 Acute Toxicity Study of Nanoparticles of Bismuth Trioxide by Inhalation Exposure in Male Rats.

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Bismuth (Bi) compounds are widely used in several products, including metallurgical and medical devices; Mexico is among the largest Bi producers. Bi toxicity is poorly studied, at micro scale it alters the kidney, nervous and reproductive systems, but little is known about the cytotoxicity of Bi nanoparticles (Bi-NP). Nanotoxicology has gained attention in the last decade due to the dramatically increase of nanomaterials in new appliances and their release to the environment. We conducted an in vivo study to evaluate the toxicity of Bi-NP by inhalation exposure. We exposed adult male Wistar rats (n=7) to an acute inhalatory dose (140 mg/kg, b.w. dispersed in 10 mg/ml BSA) of Bi2O3 nanoparticles during 5 h using the InExposure SCIREQ® inhalation system by nebulization of 1.05 mL Bi2O3 suspensions/min. The control group (n=7) received nebulized phosphate buffer in BSA. Physiological parameters such as body weight and relative organ weights, and histopathology were determined. In addition, complete blood count (CBC) and blood chemistry were evaluated. Bronchoalveolar lavage (BAL) was performed to evaluate pulmonary inflammation by differential cell count. Relative weights of spleen, liver, kidney and testes were significantly higher (12-23% increase) in the exposed group, while the histopathology revealed alterations in lungs, liver, brain and spleen, and damage in the epithelium seminiferous. CBC, blood chemistry and BAL parameters were not affected by the acute inhalation exposure to Bi-NP. In our experimental conditions, Bi2O3 nanoparticles showed moderate toxicity by inhalation exposure. Further studies are needed to evidence adverse effects due to the exposure to these NP; a subchronic study is underway to reach this objective.

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2386 Pharmacokinetics and Biodistribution of Iron Oxide Nanoparticles Using Accelerator Mass Spectrometry.

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Nanoparticles (NP) and their use as diagnostic and therapeutic agents in the biomedical field are becoming increasingly more popular. However, to date, the toxicity and biological fate have not been thoroughly investigated. Iron oxide nanoparticles are utilized for many bio-applications, which include, imaging, as drug delivery vehicles and for cell tracking. In this work, Accelerator Mass Spectrometry (AMS), an ultrasensitive technique for quantifying rare isotopes, is used to quantify the biodistribution and pharmacokinetic properties of 14C-labeled iron oxide NP in vivo.

14C-labeled carboxylated iron oxide NP (r10nm core size) were administered by nose only inhalation (0.175mg) or by a bolus intravenously (IV) (0.15mg) to male mice. For inhalation delivery, over 7 d, NP were observed to clear primarily from the lungs through the gastrointestinal system for excretion in the feces (t1/2 = 1.42 d). Detectable levels were also observed in plasma and phagocytic organs such as the liver and spleen. Furthermore, accumulation in the olfactory bulb was also observed (1.05 ng/mg at 7d). After intravenous delivery, NP were observed to accumulate primarily in the liver, spleen and lungs over a 24 h period; the half-life in plasma was t1/2 = 6.4 h. Taken together, these data indicate that once administered, carboxylated iron oxide NP are absorbed and distributed to major organs and are retained in tissue through 24 h for IV and through 7 d for inhalation. These observations may provide insight for assessment of potential toxicity upon exposure to iron oxide nanoparticles. This work performed under the auspices of the U.S. Department of Energy by Lawrence Livermore National Laboratory under Contract DE-AC52-07NA27344 and supported by LLNL CRAD No. PNNL/284.

LLNL-ABS-586092

2387 Zebrafish Xenograft Model of Glioblastoma to Identify Metal Oxide Nanoparticles with Anticancer Properties.

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Zinc oxide nanoparticles (ZnO-NPs) demonstrate selective cytotoxicity toward cancer cells in culture, and this effect may extend to other metal oxide nanoparticles (MO-NPs). Therefore, MO-NPs may possess unique qualities applicable to nanomedicine. To realize their potential as anticancer agents, we must identify safe and effective MO-NPs. We developed a screening approach that first utilizes cell culture assays to identify MO-NPs that prevent glioblastoma cell proliferation and determine the mechanism of selective toxicity. Then we assess the toxicity of the MO-NPs utilizing the embryonic zebrafish assay, an efficacious model for nano-safety assessment because of its high homology with humans and use of minimal test material. We prioritize MO-NPs that demonstrate relatively low toxicity to the zebrafish yet maintain preferential toxicity toward cancer cells in culture for assessment in a zebrafish xenograft model of glioblastoma. By xenotransplanting human glioblastoma cells into the cranium of zebrafish, we have developed an assay to identify MO-NPs that selectively inhibit cancer cell proliferation in vivo. We are testing four MO-NPs: zinc oxide, titanium dioxide, cerium dioxide, and tin dioxide. Preliminary results demonstrate that ZnO-NPs inhibit glioblastoma cell proliferation at 0.1 mM. Our screening paradigm holds promise for identifying physico-chemical traits which enhance the anti-cancer properties of MO-NPs while supporting safe NP design. Research support: NSF 134468, NIEHS P30 ES000210, ES 016896, T32 ES07060, and Air Force Research Laboratory #FA8650-05-1-5041.

2388 Toxicity of Zinc Oxide Nanoparticles: Role of Particle Dissolution and Photocatalytic ROS Production.

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Dissolution of zinc oxide nanoparticles (ZnO NPs) to ionic zinc has been recognized as an important mode of toxic action (MOA) due to their high solubility and high potency of ionic zinc to aquatic organisms. However, toxicity of ZnO NPs associated with their photocatalytic properties (generation of ROS under UV radiation) has been rarely reported. To understand the relative importance of these two MOAs, the current study investigated ZnO NP (30-50 nm) toxicity to Daphnia magna under simulated solar radiation (SSR) versus ambient laboratory light. D. magna immobilization, intracellular ROS production, and oxidative stress (lipid peroxidation) were used as toxicity endpoints. Particle dissolution was measured in a time-course manner using ICP-MS. Photocatalytic ROS production was measured by a fluorescent-based ROS assay and methylene blue photodegradation. Bulk ZnO (>1μm) and ZnCl2 were tested as reference toxicants. Concentration-dependent ROS production was detected for ZnO NPs under SSR, but not under laboratory light. Particle dissolution showed no significant differences between irradiation conditions, and both had a maximum dissolution rate of 20% within 24 h period. 48-h EC50s for D. magna immobilization under SSR and laboratory light were 0.069 mg/l (95% CI: 0.043, 0.109) and 0.109 mg/l (0.072, 0.171), respectively. Intracellular ROS production and lipid peroxidation in daphnids exposed under SSR were significantly greater than those under laboratory light, suggesting photo-induced toxicity associated with photocatalytic ROS production might have occurred. ZnCl2 had EC50s of 1.2 mg/l under SSR and 1.1 mg/l under laboratory light. Bulk ZnO showed slightly lower toxicity than ZnO NPs. Our results suggest that photo-induced toxicity of ZnO NPs related to their photocatalytic properties should not be neglected when evaluating potential environmental hazards of these nanomaterials, and the relative importance of this MOA under different exposure scenarios and towards different environmental species warrants further investigation.

2389 Luminescent Lanthanide-Doped Metal Oxide Nanoparticles for Study of Deposition and Clearance in the Respiratory Tract.


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Epidemiological studies have reported increased morbidity and mortality from respiratory and cardiovascular diseases which have close associations with exposure to airborne particulate matter (PM). Of the PM in the ambient air, ultrafine particles...
have been well-demonstrated to be one of the key pathogenic factors for cardiorespiratory disorders. Previous studies investigating deposition and clearance have used mostly radiolabeled-PM, fluorescent labeling or specially engineered nanoparticles (NPs) with physical and chemical properties that are atypical of ambient PM. A low cost and high throughput alternative is needed that will permit direct measurement of PM deposition and clearance. Here, we have used luminescent trivalent europium (Eu(III)) oxide nanoparticles (Gd2O3: Eu(III)) synthesized by a low cost cost spray flame synthesis to study the deposition and clearance of PM in rats. As rare earth elements, the lanthanides (i.e. Gd and Eu) exhibit very low natural abundance, and provide extremely good detection sensitivity in different organs with the use of inductively coupled plasma mass spectrometry (ICP-MS). Moreover, the strong optical emission that arises from the intra-4f transition of the europium ion adds a powerful tool to detect the deposition site via fluorescence microscopy. ICP-MS data from the instillation study showed that 59% of the particles remained in the lung while a significant amount was detected in the feces (20.4%) after 24 hrs, suggesting a fast clearance mechanism. Dissolution of the NPs was investigated in vitro by monitoring the photoluminescence at physiological pH and at an ionic strength for the instillation study described here.

2390 Angiogenesis Alterations Caused by TiO2 Nanoparticles.

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Angiogenesis is a process by which the preexisting vascular tree of a tissue causes growing of new blood vessels. It plays a key role in tumor development and has been demonstrated that titanium dioxide nanoparticles (TiO2 NPs) exposure increases hypoxia and growth factors levels and these events are related to angiogenesis. However, the TiO2 NPs effect in this process has been poorly explored. In this regard, chronic exposure to TiO2 NPs could develop disorders in angiogenesis. The aim of this work was to study angiogenesis alterations caused by TiO2 NPs exposure. Characterization of TiO2 NPs was done using dynamic light scattering to measure nanoparticles size distribution and zeta potential. Chicken chorioallantoic membrane (CAM) model was used as following: fertilized eggs were incubated during 7 days at 37°C and 80% humidity and then exposed to 0, 5 and 10 μg of TiO2 NPs previously suspended in MCDB-131 cell culture media and injected in blood vessels incubating for 7 days more. On 14th day, eggs were opened and digital analysis of images was done. Umblical vein endothelial cells were obtained from embryos to measure VEGF expression by flow cytometry. Results showed that suspension of 5 and 10 μg of TiO2 NPs had an agglomerated size of 535.2±93.59 nm and 503.1±1.069 nm, respectively. Zeta potential of agglomerates suspended in MCDB-131 was -17.36±1.18. Images of CAM showed structural differences between vessels of treated and untreated TiO2 NPs eggs. An increase of 0.5 cm and of 1 cm between blood vessels was found in CAM of 5 μg and 10 μg TiO2 NPs treated eggs, respectively. In addition, an increase in VEGF expression was observed in endothelial cells from TiO2 NPs treated eggs. In conclusion, TiO2 NPs exposure in CAM model induced and increased the distance between blood vessels and in the VEGF expression in endothelial cells.


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Nano scale drug delivery systems have generated considerable interest because they allow for the addition of cell specific targeting molecules and/or multiple therapeutic agents, while their bio-distribution in vivo displays molecular level kinetics. Iron based nanomaterials, with their inherent magnetic properties and an easily tailored surface chemistry, are of particular interest because of their simultaneous diagnostic and therapeutic potential. To determine how bio-distribution affects the biological efficacy of an iron based nano-carrier system, hydrophilic iron oxide nanoparticles (~10 nm) were synthesized, with either a carboxylic acid (-COOH) or an amine (-NH2) functional group. 14C labels (t1/2=5730 years) were incorporated into the organic functional groups on the surface of the nanoparticles. The radiolabeled, superparamagnetic, nanoparticles were then delivered intra-venously to mice and the pharmacokinetic distribution in vivo was determined by Accelerator Mass Spectrometry (AMS); an ultra-sensitive (10^-14 moles) quantitive spectrometric technique with small sample requirements. The radiolabeled nanoparticles were well distributed in plasma and also detected in different organs like the lungs, liver and spleen. The radiolabeling approach used in this study provides comparable bio-distribution data as the radio-labeled probes have the same chemical properties as the non-labeled probes. The radiolabeling approach described here is broadly applicable to the synthesis of nanoscale materials with multiple core and surface functionalities. The pharmacokinetic data, suggest that functionalized iron nanoparticles may have broad use as therapeutic, diagnostic or even thanerastic agents in biological systems.

PS 2392 Susceptibility to Quantum Dot-Induced Lung Inflammation Is Mouse Strain Dependent.

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Quantum dots (QDs) are nanoparticles typically composed of a CdSe core, a ZnS shell, and an assortment of polymer coatings specific to the application. Unique fluorescent and excellent semiconductor properties make QDs useful in biomedical imaging and electronics. However, due to their small size, large surface area, and heavy metal composition, there is concern over the safety of QDs. Using 8 genetically inbred mouse strains we have investigated susceptibility to QD induced lung inflammation, using % neutrophils, total protein, and levels of inflammatory cyto- tokines in bronchoalveolar lavage fluid (BALF) as biomarkers. Cadmium was measured as a marker of exposure and total glutathione (GSH) levels in frozen lung tissue were also measured. Significant treatment group and strain specific differences in the % neutrophils in BALF indicate that susceptibility to QD induced lung inflammation is mouse strain dependent. We also observed that the % neutrophils in BALF is correlated with lung Cd and GSH levels, as well as BALF cytokines. It is clear that strong relationships exist in some mouse strains and not in others. For example, the % neutrophils in BALF is highly correlated with macrophage inflamma- tory protein 1a (MIP1a) in A/J, C57BL/6j, WSB/Eij, NZO/HILj mice. However, CAST/Eij, NOD/ShiLtJ, PWK/PhJ, and 129S1/SvImJ mice do not show as strong a relationship. In future studies, recombinant inbred mouse strains will be used to map expression quantitative trait loci (eQTLs) associated with QD-induced lung inflammation. Analysis of such eQTLs could lead to insights regarding the molecular mechanisms responsible for QD toxicity and ultimately provide guidance on how to produce safer QDs. Supported by NIH grants R01ES051618, U19ES019545 and P01ES070533.

PS 2393 Nrf2 Is a Positive Regulator of Cytokine Expression in Lung of Titanium Dioxide Nanoparticles Exposed Mice.

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Background. Titanium dioxide nanoparticles (TiO2 NPs) increase the generation of reactive oxygen species and the inflammatory response in lung tissue of exposed animals. As a result, the expression of Nrf2 acts a defense mechanism against ROS generation; however its role in inflammation remains unclear. Aim. The goal of this work was to evaluate role of Nrf2 in inflammatory process induced by TiO2 NPs in lung from exposed mice. Methods. Male wild type mice (WT) and Nrf2 knockout mice (KO) were divided in the following groups: a) control-WT, b) TiO2-WT, c) control-KO and d) TiO2- KO. TiO2 NPs were suspended in 50ul of SSI and received 5 mg/kg by oropharyngeal twice a week/4 weeks. Then, mice were perfused with p-formaldehyde and lung tissue were also measured. Significant treatment group and strain specific differences in the % neutrophils in BALF indicate that susceptibility to QD induced lung inflammation is mouse strain dependent. We also observed that the % neutrophils in BALF is correlated with lung Cd and GSH levels, as well as BALF cytokines. It is clear that strong relationships exist in some mouse strains and not in others. For example, the % neutrophils in BALF is highly correlated with macrophage inflamma- tory protein 1a (MIP1a) in A/J, C57BL/6j, WSB/Eij, NZO/HILj mice. However, CAST/Eij, NOD/ShiLtJ, PWK/PhJ, and 129S1/SvImJ mice do not show as strong a relationship. In future studies, recombinant inbred mouse strains will be used to map expression quantitative trait loci (eQTLs) associated with QD-induced lung inflammation. Analysis of such eQTLs could lead to insights regarding the molecular mechanisms responsible for QD toxicity and ultimately provide guidance on how to produce safer QDs. Supported by NIH grants R01ES051618, U19ES019545 and P01ES070533.
Distinct Expression Profiles of Stress Defense and DNA Repair Genes in *Daphnia pulex* Exposed to Cadmium, Zinc and Quantum Dots.

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Use of nanocrystalline semiconductors (Quantum dots; QDs) is growing as new applications, especially in biomedical research, adopt this technology. More importantly, industrial and other mainstream uses of QDs seem likely to increase since QDs can theoretically more than double the efficiency of semiconductors in current photovoltaics. Often, with increased use comes increased appearance in the environment and, given the heavy metal composition of QDs, there exists a concern regarding the potential toxicity of these nanoparticles on aquatic organisms. The freshwater invertebrate *Daphnia* is a ubiquitous dweller of ponds and lakes throughout North America, a keystone species in aquatic food chains, and an indicator species for environmental contaminants. In this study, we aimed to compare transcriptional responses of several key stress-mediated and DNA repair genes in *D. pulex* following exposure to QDs and the individual metallic components of which they are comprised. Exposure to both Cd and QDs led to induction of mortality and Cd accumulation, which was biologically supported by the increased expression of the heavy metal responsive gene, metallothionein (MT). Our study also revealed that Cd, Zn and CdSe/ZnS QDs induced a different pattern of gene expressions regarding stress defense and DNA repair, which furthered our understanding regarding the different mechanisms of toxicity that are elicited by the nanoparticulate form of metals versus the ionic form.

A Delphi Pilot Study in Brazilian Stakeholders About Nanotechnology, Nanomaterial, and Their Toxicological and Regulatory Implications.

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With increasing numbers of nanomaterials introduced on the market in consumer applications, especially in biomedical research, adopt this technology. More importantly, industrial and other mainstream uses of QDs seem likely to increase since QDs can theoretically more than double the efficiency of semiconductors in current photovoltaics. Often, with increased use comes increased appearance in the environment and, given the heavy metal composition of QDs, there exists a concern regarding the potential toxicity of these nanoparticles on aquatic organisms. The freshwater invertebrate *Daphnia* is a ubiquitous dweller of ponds and lakes throughout North America, a keystone species in aquatic food chains, and an indicator species for environmental contaminants. In this study, we aimed to compare transcriptional responses of several key stress-mediated and DNA repair genes in *D. pulex* following exposure to QDs and the individual metallic components of which they are comprised. Exposure to both Cd and QDs led to induction of mortality and Cd accumulation, which was biologically supported by the increased expression of the heavy metal responsive gene, metallothionein (MT). Our study also revealed that Cd, Zn and CdSe/ZnS QDs induced a different pattern of gene expressions regarding stress defense and DNA repair, which furthered our understanding regarding the different mechanisms of toxicity that are elicited by the nanoparticulate form of metals versus the ionic form.

Biomarkers of Disease and Toxicity: Exploiting the Interconnections.

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With the recent focus on (a) cellular pathways involved in toxicity sequelae and (b) translational biomarkers that may link animal and *in vitro* model observations with clinical reality, there is a need to broaden the understanding of how experimental models may replicate human toxicity and disease processes. Multicellular organisms may have large numbers of genes and proteins, but a relatively limited repertoire in terms of pathophysiology in response to disease or toxicant exposure. This fact allows research to improve and protect human health to be founded on the use of model systems that allow for tractable experimentation. Animal models and, more recently, *in vitro* systems have served as a means of both exploring mechanisms and identifying hazards in terms of disease and adverse events. However, the interconnections between disease and toxicity are rarely explored leaving the information in silos. This symposium will examine organ-based toxicity and disease processes, and compare lessons learned in biomarker identification and use across toxicity, disease, and species.

In Vitro/In Vivo Assessment of Engineered Nanomaterials Using a High-Content Analysis Platform.

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With increasing numbers of nanomaterials introduced on the market in consumer products, bio-medical and environmental applications is it of primary importance to assess, understand and manage their potential toxicity. To effectively assess and understand the mechanisms of toxicity induced by nanoparticles (NPs) we implemented a platform that correlates physico-chemical properties of nanomaterials with their toxicological effects in *in vivo* on human cell lines and in *in vivo* on zebrafish as a model of toxicity to aquatic species. Nanomaterials’ physico-chemical characteristics were evaluated in the relevant exposure media for in *vitro* and in *in vivo* exposure and High Content Analysis (HCA) was employed to quantify several parameters of toxicity on exposed human cell lines and anatomical defects observed on exposed zebrafish larvae. To demonstrate the power of the platform the toxicological outcomes of surface-functionalized model nanoparticles (i.e. polystyrene Nanoparticles with carboxyl and amine surface modifications) are described along with Bismuth (Bi)-derived Nanoparticles designed for industrial applications developed within the EU/Mexico collaborative project Bionano. The model NPs were well dispersed in either complete cell culture medium or zebrafish embryo medium; for Bi-derived NPs it was necessary to develop a dispersion protocol in order to isolate particles in the nanometer range from a heterogeneous powder which were subsequently stabilized with bovine serum albumin. The HCA approach was able to differentiate between apoptosis and necrosis induced in different cell models by amine-modified polystyrene (PS-NH2) NPs and highlighted lysosomal damage as the triggering mechanism of toxicity for both the PS-NH2 NPs and the Bi-derived NPs. HCA analysis of zebrafish larvae quantified changes of anatomical features such as head/tail ratio, spine length and body curvature. The project was funded by QNano and BiNano.
Environmental contaminants and therapeutic substances contribute significantly to the high incidence and prevalence of acute kidney injury (AKI). Preclinical observations of kidney toxicity stop many compounds from entering later development due to the lack of early biomarkers of injury. In the clinic, morbidity and mortality with AKI remains unacceptably high. Our recent work has shown urinary as well as tissue fibrinogen levels increase significantly in mice, rats and humans following kidney injury/toxicity. We provide evidence that fibrinogen may serve as a key molecular link between tubulo-vascular damage and regeneration in the kidney and provides new opportunities for its use in the diagnosis and prevention of kidney disease and enablement of the clinician to institute therapeutic interventions.

Heart disease is the leading cause of death in the United States. Cardiovascular health is a prime concern for both toxicologists and physicians. Interactions among genes and environmental factors including lifestyle and drugs, may negatively affect the cardiovascular system. Myocardial damage from ischemia, infectious and metabolic diseases, or cardiotoxic drugs may be followed with biomarkers such as the troponins, natriuretic peptides, and their combination in panels with multiple evolving biomarkers. These tools facilitate an understanding of the parallels between toxicity and background disease in animal models and human heart disease.

The major goal of this symposium is to discuss the molecular and cellular mechanisms by which cytochromes P450 (CYP) contribute to oxidative stress, which could in turn lead to inflammatory processes, ultimately leading to many human diseases including cancer, neurodegenerative diseases, bronchopulmonary dysplasia (BPD), acute respiratory distress syndrome (ARDS), and drug-induced hepatotoxicity. Although much is known about the functional role of CYPs in drug metabolism, their role in endobiologic metabolism, in relation to oxidative stress and inflammation, is understudied. The recent findings of the novel role of CYPs in oxidative stress and inflammation in the manifestation of multiple human diseases warrant the need for a symposium to discuss the latest mechanistic research in this area and its impact on human health. Specifically, the symposium will discuss: i) the role of CYP4F in neuroinflammation, which in turn contributes to neurodegenerative diseases such as Parkinson's and Alzheimer's disease; ii) the role of the Arnt function as a key molecular link between tubulo-vascular damage and regeneration in the kidney and provides new opportunities for its use in the diagnosis and prevention of kidney disease and enablement of the clinician to institute therapeutic interventions.

Inflammatory processes are involved in pathogenesis and progression of CNS disorders, such as infection, traumatic brain injury, and neurodegenerative diseases. Eicosanoids including leukotrienes, particularly leukotriene B4 (LTB4) mediate inflammatory response by initiating and amplifying generation of cytokines and chemokines. Cytochrome P450 (CyP), a family of heme proteins mediate metabolism of xenobiotics and endogenous compounds, such as eicosanoids. We demonstrate that mouse brain CyPs is expressed ubiquitously in several cell types in the brain including neurons and microglia, and modulate inflammatory response triggered by lipopolysaccharide (LPS), in vivo and in microglial cells, in vitro through metabolism of LTB4 to the inactive 20-hydroxy LTB4. Chemical inhibitor or shRNA to CyP5b enhances the inflammatory response, while the PPAR agonist, fenofibrate induces CyPs and attenuates it. Fenofibrate also confers neuroprotection against Japanese encephalitis (JE), in vivo, in a mouse model of JE viral infection through up-regulation of CyP4s, and could potentially be used for prophylaxis during JE epidemics to reduce mortality and morbidity. Thus, catalytic activity of CyP4s is a novel target for modulating neuroinflammation.
the levels of 81 metabolites in several mouse organs/tissues of mice treated with TCDD. The levels of TCDD were increased in some organs but not others, suggesting that other enzymes may also be involved. TCDD increased the levels of the esterified forms of the eicosanoids in the liver in parallel with the corresponding free forms. The phospholipids so formed therefore represent a reservoir for these metabolites. Analysis of Ahr-/- null mice demonstrated that the changes in eicosanoid levels elicited by TCDD depend upon AhR. Many epoxides and diols of three to five unsaturated fatty acids, eicosapentaenoic acid (20:5n-3), docosahexaenoic acid (22:6n-3), and eicosatrienoic acid (18:3n-3), were also markedly increased in the liver, lung, but not the heart of mice treated with TCDD. Since many of the oxidized metabolites that were increased by TCDD treatment exhibit potent biological activities, including both pro- and anti-inflammatory effects, these studies lay the foundation for future experiments addressing their potential role in mediating the toxic and other effects of TCDD and other ligands of the AHR.

**2407 Regulation and Functions of Cytochromes P450 during Infection.**

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**Sponsor:** B. Moorthy.

Activation of the innate immune system, whether by infection or aseptic stimuli, causes significant changes in hepatic cytochrome P450 (P450) enzyme expression, drug metabolism and clearance. This regulation has important consequences for drug administration and responses in disease states. Its reversal by therapeutic pepti-
des targeting proinflammatory cytokines gives rise to a newly recognized drug-drug interaction mechanism. Understanding the regulatory pathways for the differ-
enst enzymes is pivotal to predicting the clinical consequences of such regulation during disease and therapy. We have studied the regulation of hepatic P450 expres-
sion in various disease models in mice, including injection of bacterial lipopolysac-
charide and infection with the colonic pathogen Citrobacter rodentium. The en-
zymes affected are dependent on the disease model, so that different drugs are likely to be affected in different human disease states. In the C. rodentium-infected mice, increases in cytokine or cytokine receptor-null mice indicate roles for interleukin-6 and tumor necrosis factor-α in the regulation of a small subset of P450 transcripts, in-
cluding some members of the Cyp3a subfamily. Studies in SCID mice suggest that T cells or T-cell-derived cytokines might also be important in this model. However, the absence of cytokine signals fails to alter the down-regulation of the most pro-
foudly suppressed liver enzymes i.e. Cyp4a10, Cyp4a14 and flavin monooxygenase 3 (Fmo3), suggesting a different mechanism of regulation for these genes. To ad-
dress whether or not Cyp4a enzymes play a role in the host-pathogen interaction, we infected Cyp4a14-/- and Cyp4a10-/- mice with C. rodentium, and compared their responses to those of wild type mice. Results from these ongoing studies will be presented. Supported by National Institutes of Health grant R01DK072372.

**2408 Mechanistic Role(s) of Cytochrome P450A1 and B1 Enzymes in Hyperoxic Lung Injury: Implications for Bronchopulmonary Dysplasia (BPD) in Premature Infants and ARDS in Adults.**

B. Moorthy. *Pediatrics, Baylor College of Medicine, Houston, TX.*

Hyperoxia is routinely used in the treatment of pulmonary insufficiency and respi-
ratory distress in preterm and term infants and in adults with acute respiratory dis-
ease (ARDS). However, in premature infants, hyperoxia contributes to the de-
velopment of chronic lung disease (CLD), which is termed bronchopulmonary dysplas-
ia (BPD). The molecular mechanisms of oxygen-mediated lung injury are not un-
derstood, but reactive oxygen species (ROS) are the most likely candidates. ROS-me-
diated reactions with biological macromolecules such as DNA, proteins, and lipids are also responsible for many other lung diseases such as acute respiratory distress syndrome (ARDS), asthma, emphysema, chronic obstructive pulmonary disease (COPD), and lung cancer induced by environmental pollutants. Results from our laboratory demonstrate a novel role for cytochrome P450 (CYP1A) enzymes in the detoxification of ROS-mediated lipid peroxidation products, e.g., F2-isoprostanes. Our major observations are that mice lacking the genes for Cyp1a1 or 1a2 are more susceptible to hyperoxic lung injury than wild type mice, with Cyp1a2-null mice being the most sensitive. On the other hand, mice lacking the gene for Cyp1b1, are less susceptible to lung injury, suggesting a pro-oxidant role for Cyp1b1. Mice pre-treated with the CYP1A inducer β-naphthoflavone (BNF), fol-
lowed by exposure to hyperoxia leads to protection against lung injury. We also found formation of bulky oxidative lesions (oxidative DNA adducts) in tracheal as-
pirates of premature infants and adults who received supplemental oxygen, and this was associated with BPD and ARDS, thereby suggesting that these adducts could serve as novel biomarkers of these diseases. Future studies could lead to the de-
velopment of rational strategies for the prevention/treatment of lung diseases associ-
ated with hyperoxia.

**2409 Molecular Basis of Age-Related Susceptibility to Chemicals and Environmental Hazards: From Model Systems to Humans.**

J. S. Lee1 and J. C. Fuscoe2. *US EPA, Durham, NC; US FDA, Jefferson, AR.*

The susceptibility of individuals to chemicals and environmental hazards at the ex-
tremes of the population age-distribution (the very young and the very old) is often not adequately assessed. By understanding genes expressed at the various life stages, the assessment of health risk versus benefit can be more rationally determined. Children are more susceptible to exposures to environmental toxicants compared to adults because intakes are increased, biologically-effective doses may differ, and early lifestyle exposure may lead to adverse health effects that are chronic in nature. Older adults are more susceptible to environmental toxicants because of pharmaco-
kinetic and pharmacodynamic changes associated with aging. Altered absorption, distribution, metabolism, and excretion (ADME), along with decreased blood flow to the liver, decreased liver mass, and decreased content of specific cytochrome P450 (CYP) could result in decreased clearance of chemicals in older adults. In addition to these factors, genetic and epigenetic changes that occur with age and chemical exposure may also increase susceptibility to environmental hazards. Our panel will examine genomic and epigenomic changes that occur with age using animal and human data. Rat liver and kidney data will be used to discuss age-related target organ vulnerabilities. Human relevance will be addressed using a longitudinal birth cohort study in Mexican-American children, two mother-child cohort studies in Mexico and the United States, and a cohort study of older men from the Normative Aging Study. Ultimately, understanding the implications of genetic and epigenetic changes related to age on the effects of chemical exposure will help to protect the health of children and older adults. Disclaimer: The views expressed are those of the authors and do not necessarily represent the views or policies of the US EPA or US FDA.

**2410 Functional Genomic and Epigenomic Changes in the Liver and Kidney during the Rat Life Cycle.**

J. C. Fuscoe, T. Han, V. Vijay, V. Desai and J. C. Kwekel. *Division of Systems Biology, NCTR, US FDA, Jefferson, AR.*

The susceptibility of individuals at the extremes of the population age-distribution (the very young and the very old) and differences between sexes are often not ade-
quately assessed. By understanding the genes expressed in each sex at the various life stages, the assessment of health risk versus benefit can be more rationally deter-
mined. Comprehensive analysis of the transcriptome, including miRNA, and the epigenome in the liver and kidney of Fisher 348 rats from 2 weeks to 2 years of age reveals substantial differences at various life-stages and between the sexes. In the liver and kidney, the expression of nearly 4000 genes was found to significantly vary with age and/or sex. Many of these genes are involved in xenobiotic metabolism and transport, processes that impact drug efficacy and safety. The expression of genes that code for some of the key cytochrome P450 biomarkers recently qualified by the FDA were found to vary substantially with age and sex, in some cases up to 100-fold. Such dramatic differences may impact the interpretation of biomarker re-
examination of miRNA expression in the liver showed nearly 200 miRNAs varied with age and/or sex. Notably, a group of 42 miRNAs was expressed at a rel-
tively high level at 2 weeks of age in both sexes followed by low or no detectable ex-
pression at older ages. However, this 2 week-specific miRNA expression was not evi-
dent in the kidney. Analysis of potential target miRNAs for liver miRNAs suggests roles in disease susceptibility involving fibrosis as well as regulation of xenobiotic metabolism related genes. A similar number of kidney miRNAs also displayed sex and age-related changes which may be linked to gene regulation. These differences may be related to age- and sex-specific susceptibilities to adverse drug reactions or disease states. Understanding these differences should improve personalized medi-
cine both in terms of disease prevention and management, and safer use of drugs.

**2411 A Case-by-Case Approach to Pediatric Drug Safety Involving Multiage Juvenile Rat Models That Target Developmental Issues and Address Regulatory Concerns.**

P. Espandiari, W. Rodriguez and J. P. Hainz. *CDER, US FDA, Silver Spring, MD.*

The toxicity of drugs in pediatric populations may vary considerably from that seen in adults. Capacity to generate or inactivate the toxic moiety, issues of drug half-life, volume of distribution or specific target organ toxicity as well as hypersensitivity are
all critical factors. Vital organ systems mature at different times and present many challenges for selecting an appropriate model that captures the vulnerability necessary to characterize safety issues. One example of a multi-age pediatric model utilizes rats 10, 25, 40 and 80 days old roughly corresponding to human infants, toddlers, teenagers and young adults exposed to either the nephrotoxins gentamicin, or cisplatin or the hepatotoxic valproic acid. The differential effects across the various age groups were assessed by organograms, histopathology, hematology, clinical chemistry, liver and kidney biomarkers, and metabolomics technologies to establish specific age-related target organ vulnerabilities. Each age group was shown, using principal component analysis, to be a distinctly separate entity in terms of treatment responses. This published model, representing a single possible approach, has potential for significant utility as a tool for pediatric safety evaluation, in comparison to dosing throughout the complete maturation process. The FDA Guidance for Nonclinical Safety Evaluation of Pediatric Drug Products (Feb., 2006) presents a wide variety of parameters involving juvenile/adult differences. These constitute many possible choices that may be incorporated into a pediatric preclinical model appropriate to address preclinical safety. The age of intended exposure, route, frequency and length of exposure are critical and will be discussed. In general the preclinical safety evaluation of new and approved drugs for pediatric use is handled on a case by case basis. This is reflected in some of the more recent EMA and ICH documents (2008, 2009) that allow for a multitude of choices based on the ultimate requirements of subsequent clinical trials.

**2412 Genetic and Epigenetic Mechanisms of Susceptibility to Environmental Exposures in Children.**


Children are more susceptible to exposures to environmental toxicants than adults, and in utero exposures may result in developmental problems and chronic diseases. Some children can be particularly vulnerable due to their genetic makeup and age-related differences in protective enzyme levels such as paraoxonase (PON1). Results from the Center for the Health Assessment of Mothers and Children of Salinas (CHAMACOS) birth cohort study will be presented to illustrate complex relationships of the effects of prenatal exposure to pesticides, PON1 functional genomics, and genome-wide DNA methylation in Mexican-American children. We determined PON1 genotypes and three PON1 enzyme activities in 450 children and their children. Although it was previously thought that children’s PON1 levels reach adult ones by age 2, we found that PON1 levels and activities were the lowest in newborns and steadily increased with age. However, they remained below adult levels up to 7 years. Infants and young children, particularly those with PON1 genotypes encoding for lower PON1 levels and activities, have up to 65-fold lower levels of the protective PON1 enzyme than adults and may be especially susceptible to OP exposures. To assess DNA methylation in cord blood and peripheral blood (clots) of 9 year old CHAMACOS children we interrogated 485,577 CpG sites from >24,000 genes using the Illumina BeadChip platform. We found that 15.5% of all CpG sites were differentially methylated between children at birth and 9 years of age. More than 2% of CpG sites investigated, in >1,900 genes, showed significant differences in methylation by sex, including 731 CpG sites located in autosome differentially expressed and pathways involved in response to environmental exposures during pregnancy have been identified. Unlike genes, epigenetic mechanisms could be reversible and an enhanced understanding of their role may lead to better protection of pregnant women and children, and improved public health.

**2413 Prenatal Metal Exposure and Health Effects.**

B. C. Fry. Environmental Sciences and Engineering, University of North Carolina at Chapel Hill, Chapel Hill, NC.

We will present data that we are collecting from two mother-child cohorts, one in Gómez Palacio, Mexico and one in North Carolina, USA. Both of the newborn cohorts are at risk for metal exposure. The research focuses on an arsenic endemic area of Mexico, Gómez Palacio, the study site for our newly established birth cohort. In this region we are assessing exposure to inorganic arsenic in pregnant women. Our results suggest that pregnant women in this region are at risk for high level exposure to arsenic, a finding that has implications both for the health of the women and also their children. In related work we have discovered some aspects of arsenic are associated with changes in DNA methylation patterns. We are also examining changes in DNA methylation patterns in fetal DNA collected from the NC cohort which is at risk for prenatal exposure to cadmium. We demonstrate using genome-wide gene-specific DNA methylation analysis that there are significant differences in levels of promoter methylation in fetal or maternal samples. In addition, we highlight that prenatal exposure to cadmium is associated with altered patterns of methylation of genes in the placenta.

**2414 Particulate Pollution, Susceptibility, and Epigenetic Pathways in an Elderly Cohort.**

J. Madrigano1, A. Baccarelli2, M. A. Mittleman1, R. O. Wright2, D. Sparrow3, P. S. Vokonas1, L. Tarantini1 and J. Schwartz2. 1Mailman School of Public Health, Columbia University New York, NY. 2Autonomic Laboratory of Public Health, Boston, MA. 3Beth Israel Deaconess Medical Center, Boston, MA. Sponsor: L. Lee.

DNA methylation is a potential pathway linking environmental exposures to disease, and lower DNA methylation has been found in processes related to cardiovascular morbidity. Genetic and other host characteristics, such as psychological functioning, have been found to modify the association between air pollution and morbidity. In our study, DNA methylation of repetitive elements, as well as specific genes, was measured in 1406 blood samples from 706 elderly participants in the Normative Aging Study (NAS). We will discuss the changes in repetitive element DNA methylation, as well as two specific genes (the inducible nitric oxide synthase gene, iNOS, and the glucocorticoid receptor gene, GCR) associated with ambient particles (PM2.5) and black carbon (BC), estimated with mixed models. We will also discuss genotype and phenotype characteristic that may modify this association.

**2415 Challenging the Limits of Nonclinical Safety Assessment of Pediatric Medicines.**


Pediatric safety assessments are a fundamental and integral part of drug development programs. Introduction of regulatory guidance in the last decade has formalized the inclusion of safety evaluations in juvenile animals, leading to a better understanding of potential drug effects on developmental processes and risks specific to pediatric age groups. Toxicology studies in juvenile animals have evolved to address inherent differences in susceptibility between mature and immature systems. Pediatric safety assessments are challenged by practical and interpretive complexities of conducting toxicity studies in immature animals against a background of increasingly diverse disease indications. This symposium will review a number of innovative approaches that challenge the current limits of the nonclinical safety assessment of new pharmaceuticals. Our panel of experts will discuss unique approaches and case studies dealing with the challenges of supporting pediatric formulations, nontraditional routes of administration, and the complexities of developmental neurotoxicity assessments. Furthermore, our experts will discuss how information from various sources such as in vitro experiments using neonatal tissue, pharmacokinetics, and clinical pharmacology work may be brought together to build a risk assessment specific for a young infant and pediatric dosing. These innovative approaches from industry and government challenge the limits of nonclinical safety assessment and provide reassurances of safety for pediatric medicines.

**2416 Safe and Effective Use of Established and Novel Excipients in the Development of Children’s Medicines.**


Developing medicines for children is in some respects even more challenging than developing medicines for adults yet the range of excipients available for use in the development of children’s medicines is likely to be significantly reduced without appropriate evidence of safe, tolerated excipient limits across the whole paediatric age range. Industry, with the support of academia, have the opportunity to work with
national regulatory agencies to raise awareness of the need to build a risk/benefit approach focused on the use of excipients in paediatric drug products; in particular, the use of novel excipients in this population. Identification of key data should be determined together with an appropriate mechanism for this data to be shared to enhance access to medicines for children.

**2417 Technical Challenges and Data Interpretation Evaluating an Inhaled Long Acting β2-Agonist in Juvenile Dogs.**

A. Mous. Nonclinical Drug Safety Germany, Boehringer Ingelheim Pharma GmbH & Co. KG, Biberach an der Riß, Germany.

Numerous inhaled medications are marketed to pediatric populations, but most of these products were registered prior to the regulated integration of juvenile toxicity studies in nonclinical safety programs. In order to initiate clinical trials in pediatric populations, safety assessments are conducted in juvenile animals to assess any potential toxicity effects during postnatal development, particularly with regard to lung development. This presentation will detail a case study for an inhaled, long acting β2-agonist in juvenile dogs. Discussion will include insight into the pediatric safety strategy, technical challenges involved in dosing an inhaled medication to juvenile dogs, and data interpretation.

**2418 Modeling Pediatric Exposures to Neuroactive Agents: Developmental Neurotoxicology.**

M. G. Paule. Division of Neurotoxicology, National Center for Toxicological Research, US FDA, Jefferson, AR.

Given the obvious limitations, it is difficult to thoroughly explore the effects of pediatric neuroactive agents on neurons in human infants or children. Due to the complexity of the primate brain, the monkey is often the animal model of choice for developmental neurotoxicology experiments. Case studies will be presented on nonhuman primate models of pediatric exposures to general anesthetics and other neuroactive compounds. These examples will focus on translational endpoints involving cognitive functions such as learning and memory and in vivo imaging using PET. Strategies for neuroprotection will also be discussed.

**2419 Technical and Scientific Challenges in Risk Assessments for Neonates and Infants.**


Progression into clinical trials in neonates and infants often involves a risk assessment specific for that age group. The risk assessment is likely to include information from technically challenging bespoke studies in very young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals but may also include in vitro experiments using neonatal tissue (exploring these studies may not be limited to just repeat dose toxicity studies in young animals). However, the risk assessment is likely to include information from technically challenging bespoke studies in very young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animals. However, these studies may not be limited to just repeat dose toxicity studies in young animal...
identifying QSARs that are applicable within each group. Due to high variability in the molecular structure and different mechanisms of action, individual groups of nanoparticles should be modeled separately. In each case, according to the general QSAR rules, the applicability domain of the models should be carefully validated. Our recent ab initio study revealed details of interactions of gold clusters, carbon nanotubes and fullerenes with DNA bases and base pairs. Direct prediction of toxicity of unknown nanomaterials was done using QSAR models developed for a test set of compounds characterized experimentally. Based on experimental testing, we developed and tested novel interpretative nano-QSAR model describing cytotoxicity of 17 nano-sized metal oxides to bacteria Escherichia coli. The proposed model allowed us to formulate a hypothesis that mechanistically explained differences in toxicity between the individual oxides.

2424 Biological Surface Adsorption Index (BSAI): A Molecular Signature for Nanomaterial Interactions.
J. R. Riviere, Institute of Computational and Comparative Medicine, Kansas State University, Manhattan, KS.
Most characterization techniques available today for nanomaterials are based on hard physical-chemical properties of size, shape, and surface properties, often determined under very non-biological conditions. However, a major factor that determines biological interactions of nanomaterials in vivo are their surface properties related to forming interactions with molecules in the biological environment. We have developed the BSAI metric, which develops a signature of nanomaterial surface properties specifically related to biomolecular interactions. These properties are developed based on how a nanomaterial interacts with a series of probe compounds using a QSAR approach to generate five molecular descriptors that could be described as a multidimensional partition coefficient. This presentation will introduce the index and illustrate how it can be used to improve modeling of biological interactions.

2425 Predictive Modeling on Nanoparticle-Biomolecular Interactions.
R. Pandey, Department of Physics, Michigan Technological University, Houghton, MI. Sponsor: S. Hussain.
Nano-scale materials, such as semiconductor and metal quantum dots, carbon nanotubes, and graphene, exhibit novel optical, electrical, and magnetic properties that can be exploited for new generations of electronics and sensors. Nanoscale materials also exhibit unique affinity with biological molecules, such as nucleic acids and proteins, which can be utilized for a wide variety of biological diagnostics and sensing applications. In order to develop such application concepts, however, a fundamental understanding of the interactions between various nano and biological systems is critically important. This talk will present a brief overview of recent developments in the assembly and structure-property characterizations of hybrid nano-bio materials with a focus on the physical and chemical properties of their interface. Results obtained from recent theoretical and experimental investigations on optical protein-QD and nucleic acid base-nanotube (C, BN) interactions will be presented.

2426 Toxicogenomics in Risk and Safety Assessment: Recent Advances and Continuing Challenges.
C. Thompson and M. D. Waters, 1TanStrategies, Inc., Katy, TX; 2Integrated Laboratory Systems, Inc., Research Triangle Park, NC.
Toxicogenomic studies can provide a vast amount of data with regard to the changes a chemical can have on a cell, tissue, or organism, and technological achievements continue to make it easier and cheaper to generate such data. However, the application of toxicogenomic data to environmental risk assessments and pharmaceutical safety assessments has progressed more slowly and, despite recent advances, challenges remain as to how best to harvest and interpret these large and complex datasets to facilitate their practical application. This session will describe recent applications of toxicogenomics in environmental risk assessment with a focus on assessing and predicting genotoxic modes of action and utilizing transcriptome changes from multidose and multi-endpoint animal bioassays in quantitative risk assessment. In addition, recent advances in the usage of toxicogenomics in preclinical pharmaceutical safety, clinical trial placement, and individualized medicine will be described.

2427 Characterizing and Predicting Modes of Action of Carcinogenicity Based on Conventional and Toxicogenomics Methods.
M. D. Waters, Integrated Laboratory Systems, Inc., Research Triangle Park, NC.
Predictive toxicogenomics uses global molecular expression data resulting from genomic perturbation (e.g., transcript profiles) to predict a toxicological outcome, such as carcinogenicity. In the context of risk assessment, the classification of carcinogens as genotoxic or nongenotoxic has become an essential and debatable issue because of the default assumption that drives regulatory decision-making regarding the presumed linearity of the dose-response curve for genotoxic carcinogens. In fact, the great majority of known human carcinogens are easily detected in conventional short-term tests for genotoxicity and induce tumors at multiple sites in rodents, thus provoking challenges as to the human relevance of nongenotoxic rodent carcinogens. Toxicogenomics studies appear quite useful in resolving this dichotomy and in pursuing mechanisms of action. In toxicogenomics studies, a strong DNA damage response at the gene expression level suggests direct DNA modification whereas increased expression of genes involved in cell cycle progress is more characteristic of the indirect-acting agents such as those that induce oxidative stress. Gene expression profiles have been demonstrated that discriminate nongenotoxic modes of action (e.g., cytotoxicity and regenerative proliferation, xenobiotic receptor agonists, peroxisome proliferator-activated receptors, or hormonal-mediated processes) and other profiles appear to delineate various paths to the formation of conventional cytogenetic alterations. The evidence accumulated to date suggests that toxicogenomics approaches will be useful in conjunction with conventional test methods in the dose and phenotype anchored assessment of chemical carcinogenicity. Case studies will be used to illustrate these points.

2428 Case Studies of Dose-Response Genotoxicity and Toxicogenomic Studies Designed to Replace Default Assumptions Used in Carcinogenic Risk Assessments.
L. Recio, Integrated Laboratory Systems, Inc., Research Triangle Park, NC.
Characterizing dose-response is a fundamental aspect of toxicology and can be used to determine and predict the potential adverse effects of chemicals to humans. Only recent genetic toxicity and toxicogenomic studies have adequately characterized dose–response over a range of exposures and have used these data for point-of-departure (POD) calculations needed in risk assessment, such as benchmark dose 10 (BMD10). Genotoxicity and toxicogenomic endpoints can be considered as biomarkers of ‘key events’ or adaptive responses that with robust experimental designs can provide the needed qualitative and quantitative dose-response information to establish chemical-specific modes-of-action that can be integrated into weight-of-evidence-based approaches for risk assessments. More recently genomic signatures for mode-of-action (MOA) (e.g., genotoxic vs nongenotoxic MOA) in target organs are emerging as mRNA biomarkers of effects. Low dose studies designed to identify exposure levels that do not cause alterations in basal genotoxicity or gene expression can be used to identify exposure levels (or dose) that represent the transition between the NOEL concentrations and other PODs such as BMD. Recent studies conducted at ILS with collaborators, represent case studies aimed at identifying the NOEL concentrations and other PODs using the in vitro micronucleus and mutagenicity studies in human cells for acetalddehyde, dose-response and impact of liver GSH detoxication on naphthalene genotoxicity, expression profiling studies to assess genotoxic vs nongenotoxic MOA in human TK6 cells, and an in vivo dose-response study conducted with the mouse liver carcinoma furan examining impact on the mouse genome and epigenome.

2429 Application of Transcriptomic Data for Quantitative Chemical Risk Assessment.
R. S. Thomas, The Hamner Institutes for Health Sciences, Research Triangle Park, NC.
Current challenges facing chemical risk assessment are the time and resources required to meet the data standards necessary for a published assessment and the incorporation of modern molecular biology information. The integration of transcriptomic data into regulatory decision-making may address some of these challenges by providing an efficient means to quantitatively and comprehensively evaluating the molecular changes resulting from chemical exposure. To assess the value of applying transcriptomics in quantitative chemical risk assessment, a series of studies was performed. In the first study, mice were exposed for 13 weeks to multiple concentrations of six chemicals that were positive in a two-year cancer bioassay. In a second study, mice were exposed in a time course to multiple concentrations of six chemicals with published risk assessments. In both studies, histological changes were evaluated and transcriptional microarray analysis was performed on the target tissues.
The histological and the tumor responses were analyzed using standard benchmark dose (BMD) methods to identify non-cancer points-of-departure. The dose-related changes in gene expression were also analyzed using a BMD approach and grouped based on signaling pathways. The transcriptional BMD values showed a high degree of correlation with apical responses for specific pathways and many of the correlated pathways have been implicated in relevant disease pathogenesis. Importantly, transcriptional points-of-departure for even the most sensitive pathway were on average less than three-fold different than traditional apical points-of-departure for both cancer and non-cancer endpoints suggesting that transcriptomic changes in signaling pathways can be used to estimate noncancer and cancer points-of-departure for use in quantitative risk assessments.

2430 Challenges and Opportunities of Toxicogenomics Analyses in Safety Assessment during Preclinical Safety Studies.
C. Karbowsk. Discovery Toxicology, Amgen, Thousand Oaks, CA.

Microarray analysis is a key tool utilized in the biotechnology/pharmaceutical industry as part of a holistic approach to predicting and understanding mechanisms of toxicity of molecules in development. Historically, researchers have faced significant challenges in analyzing and interpreting these large datasets such as understanding the translation of molecular changes to phenotypic changes in an organism and the relevance of observed alterations to other species. However, progress in overcoming these challenges continues to be made. For example, contextualization of gene expression changes utilizing historical and publically available reference toxicant datasets coupled with recent advances in Systems Biology such as more comprehensive and toxicologically relevant content along with the incorporation of transcription factor analyses and gene directionality provides opportunities to understand the genesis and cross-species relevance of pre-clinical phenotypic changes. This presentation will provide an historical view of the utility of microarray analysis in preclinical safety assessment and then illustrate, through case examples, the putative biological assertions. The platform was initially applied to investigative toxicity safety assessment and then illustrated, through case examples, the added value provided when current state of the art tools are applied to help dissect molecular changes underlying phenotypic alterations.

2431 Role of Causal Reasoning in Patient Stratification from Efficacy and Safety Perspectives.
A. Enayettallah. Drug Safety Research & Development, Pfizer, Inc., Groton, CT.

Advances in genomic technologies have led to the ability to rapidly generate extraordinary amounts of data. However, a lack of efficient tools to manage and integrate such large amounts of data has limited the application of genomics in the pharmaceutical industry. Recently, we developed a computational platform we call the Causal Reasoning Engine (CRE) that is a powerful tool fortranscriptomic data analysis. The CRE provides explanation of the observed transcriptomic changes in the context of prior biological knowledge, captured in a knowledge base of computable biological assertions. The platform was initially applied to investigational toxicology to provide mechanistic understanding of organ toxicities, including drug-induced liver injury (DILI) and drug-induced cardiac injury. More recent developments of the CRE platform indicate potential predictive power for evaluating compound properties to identify liabilities at the individual patient level, which would enable the stratification of patients in clinical trials. In the context of efficacy we will show a use case for patients with diseases known for their heterogeneity, such as systemic lupus erythematosus and inflammatory bowel disease. In this case individualized analysis using CRE clearly classifies patients based on their underlying disease mechanisms, and we will discuss the potential impact on patient and treatment selection in clinical trials. Finally, the results from the individualized analysis approach in patients with immune-mediated DILI will also be presented. Based on the examples presented here, we believe that the CRE approach shows great promise in being able to stratify patients to support the development of more effective and safer medicines.

2432 Assessment of Environmental, Dietary, and Biological Risk Factors Impacting Liver Cancer Incidence in Texas.
E. D. Bruce,1, 2 and A. Romoser1. 1Institute of Biomedical Sciences, Baylor University, Waco, TX; 2Institute of Ecological, Earth, and Environmental Science, Baylor University, Waco, TX.

Liver cancer is the 12th most commonly diagnosed cancer in the US and the 6th most common cause of cancer deaths. Liver cancer incidence has been increasing in both Texas and the US. Although liver cancer only accounts for 1.3% of new cancer cases, it accounts for 2.6% of cancer deaths. Survival rates are poor with a five-year survival of 13 to 15 percent. In 2012, approximately 2,197 Texans are expected to be diagnosed with liver cancer and 1,768 are expected to die from the disease. Liver cancer incidence trends for Texas and the US were determined using data from the Department of State Health Services, Texas Cancer Registry and National Cancer Institute, Surveillance Epidemiology & End Results. For a 15-year period (1995 to 2009) age-adjusted incidence rates were computed by gender and race/ethnicity. Texans experience higher liver cancer incidence rates than the US and the rates are increasing faster. From 1995 to 2009 Texas rates increased by an average annual rate of 5.7% compared to 3.9% for the US. The 15-year percent changes in incidence for Texas and the US were 126% and 77%, respectively. In Texas, liver cancer incidence has been increasing in both men and women but faster in men. From 1995 to 2009 age-adjusted rates for men increased by 130%, while rates for women increased by 96%. In both Texas and the US, men of all race/ethnic groups are diagnosed two to four times as often as women. In Texas, Hispanics (any race) and Asian/Pacific Islanders have the highest rates but Blacks had the highest rate of change from 1995 forward with an annual percent increase of almost 7% and a 15-year increase of 141%. Texas Hispanics and Blacks have significantly higher liver cancer incidence rates than US Hispanics and Blacks.

2433 A Lay Health Worker-Based Intervention for Reducing Families’ Environmental Exposures.
L. Cizma1, J. Ross1, R. Rincon2, H. Tamez2, A. Ginez2, R. Perale2, C. Miller2 and T. McDonald1. 1Texas A&M Health Science Center School of Rural Public Health, Texas A&M University, College Station, TX; 2South Texas Environmental Education and Research Program, University of Texas Health Science Center, San Antonio, San Antonio, TX.

In economically disadvantaged areas of San Antonio and Laredo, TX, poor living conditions and a hot climate increase the likelihood of pest infestations, leading to increased pesticide use. Promotora (lay health workers) were employed to deliver a pesticide health education module to families with children between 6 months to 5 years of age. Assessments of attitudes and behaviors relating to pesticide use were given prior to and 6 months after module delivery. In Laredo, participants reported statistically significant changes in behavior and attitudes six months after the module. For example, the percent of participants reporting that they always had emergency numbers by the phone was 35% before the training and 86% six months after.
after the training, and the percentage reporting that they always used gloves when applying pesticides was 35% before the training and 84% six months after the training. Promotora-driven health education with local collaboration may improve family attitudes and practices relating to pesticide use, and may be explored as an option for addressing other environmental exposures as well.

Despite the public health problem of obesity and increasing incidence of hepatocellular carcinoma (HCC) in the United States (US), the relationship between obesity and HCC has never been examined extensively in US population. At the University of Texas MD Anderson Cancer Center we conducted a case-control study aimed at examining HCC risk factors in the US. Cases were patients with pathologically confirmed diagnosis of HCC and US residency. The healthy control subjects were spouses of patients at MD Anderson who had cancers other than liver, gastrointestinal, lung, or head/neck cancer. Self-reported weight and body size (Stunkard pictograms) at ages 20, 30, 40, 50, 60, 70 was obtained from participants by personal interview. Between 2005 and 2011 we enrolled 403 cases and 661 controls. Body mass index (BMI) was classified as "underweight" (BMI < 18.5), "normal" (BMI range, 18.5-24.9), "overweight" (BMI range, 25-29.9), and "obese" (BMI ≥ 30.0). We found that individuals who were obese from the ages of 30 to 49 had a significantly increased risk of HCC, independent of HCC established risk factors. The estimated odds ratio (OR) and 95% confidence interval (CI) was 4.1(1.6-10.5). The association was observed in men and women; the ORs (95% CIs) were 2.3(1.1-4.9) and 2.9(1.2-8.9) respectively. Moreover, individuals who were overweight or obese from the ages of 30 to 49 years had an earlier onset of HCC by 3 to 6 years (median age of onset was 65 years for patients with normal weight, 62 years for overweight patients [P=0.02], and 59 years for obese patients [P <0.001]). Underlying evidence of cirrhosis was significantly observed in HCC patients with early obesity. We concluded that obesity is a significant risk factor for HCC in the US where underlying cirrhosis can be a significant burden in disease management. Integration of obesity with other HCC risk factors into a risk model may lead to the development of a new scoring system to identify high-risk individuals who may benefit from HCC screening and prevention.

Biomarkers of Hepatocellular Cancer Risk and Diagnosis.

R. M. Santella, NIEHS Center for Environmental Health, Columbia University Mailman School of Public Health, New York, NY.

Hepatocellular carcinoma (HCC) incidence is increasing in the US and HCC has one of the fastest growing death rates of any cancer. There is wide geographic variation in HCC incidence around the world likely due to geographic differences in the prevalence of various etiological factors. In Asia and Africa hepatitis B virus is the primary etiologic agent while in the US, hepatitis C virus is more common.

Identified environmental/lifestyle risk factors include aflatoxin B1 (AFB1), a dietary mold contaminant, alcohol drinking, and cigarette smoking. We have used biomarkers in a prospective study in Taiwan to demonstrate that elevated baseline levels of AFB1 urinary metabolites, AFB1-albumin adducts, polycyclic aromatic hydrocarbon (PAH)-albumin adducts and urinary isoprostanes, a biomarker of oxidative stress, are associated with later development of HCC. We have also found AFB1- and PAH-DNA adducts in liver tissues of US HCC cases. Using a candidate gene approach, studies of liver tumor tissues have identified genes that are hypermethylated in tumors compared to adjacent tissues and found that this methylation was associated with elevated levels of AFB1-DNA adducts. DNA isolated from plasma collected at the time of diagnosis contains these same methylated markers. More importantly, methylated DNA released from the tumor can be frequently found in plasma collected many years before clinical diagnosis suggesting their potential utility in screening high populations such as those with viral infection. More recently, we have used Illumina Infinium arrays that interrogate methylation of 27k or 450k CpG sites to more comprehensively identify regions with altered DNA methylation. These studies have found large numbers of hyper or hypomethylated regions in tumors compared to adjacent tissues and identified methylation markers that should enhance early diagnosis. Biomarkers of environmental exposure in combination with viral infection markers can identify populations at increased risk while methylation and microRNA markers may potentially be used for the early diagnosis of HCC.

Mitigation of Aflatoxin Exposures Using a Clay-Based Enterosorbent.

T. D. Phillips, Veterinary Integrative Biosciences, Texas A&M University, College Station, TX.

Concerns about the quality and safety of foods destined for human and animal consumption have evoked a growing awareness of the significant hazards associated with chemicals known as aflatoxins. The aflatoxin problem in foods is longstanding, unavoidable and seemingly inextricable. Aflatoxin exposure is often considered a risk factor for disease in countries where there is a lack of infrastructure for food safety regulation. However, aflatoxin exposure has been observed and is a cause for concern in underprivileged communities in Texas and the Southwest US as well.

Aflatoxin B1 (AFB1) is a direct acting mutagen and has been shown to disrupt genes involved in carcinogenesis and tumor suppression. Recent research with mycotoxin enterosorbent, NovaSil (NS), in an African population has shown the product to be safe and efficacious in reducing biomarkers of exposure to aflatoxin. Blood and urine samples were taken at Baseline, 1 Month, 3 Months and 4 Months. NS clay significantly reduced biomarkers for aflatoxin exposure in urine and blood and the treatment was well tolerated by participants. In a recent San Antonio study focusing on a population with a significantly elevated incidence of liver cancer, it was determined that biomarkers for aflatoxin correlated with ingestion of foods known to contain relatively higher levels of aflatoxin. Phase II of this study is underway to determine NS efficacy in reducing exposures to this toxin in Texas. Mitigating AFB1 exposure using NS represents an innovative, practical, sustainable and environmentally benign approach that will benefit more than 4.5 billion people living in climates conducive to the growth of fungi and production of mycotoxins in staple foods.

Risk Factors Influencing the Incidence of Liver Cancer in San Antonio.

F. A. Guerra, Department of Pediatrics, University of Texas Health Science Center San Antonio, San Antonio, TX. Sponsor: E. Bruce.

The incidence of hepatocellular carcinoma (HCC) is significantly elevated in Hispanic communities in Bexar County, Texas. Multiple factors including diet, environment, occupation, lifestyle, health status, and gender play a role in the etiology of HCC. Previous research with Texas A&M University has focused on defining the risk factors that may influence the high incidence of liver disease observed in San Antonio. Epidemiological and clinical intervention studies are ongoing in Bexar County, and have been successful in raising awareness in the community regarding environmental, dietary, and biological risk factors for disease. Ongoing studies with The University of Texas Health Science Center-San Antonio and Texas A&M University will investigate the impact of an intervention trial designed to decrease biomarkers of mycotoxin exposure and enhance public health in communities at high risk for HCC.

A Novel Single Cell-Based High-Throughput Toxicity Study of Drugs.

L. Ma, Y. Qiao and M. Su, NanoScience Technology Center, University of Central Florida, Orlando, FL. Sponsor: T. Lam.

Many anticancer drugs are genotoxic. The ability of tumor cells to repair drug induced DNA damage is indicative of therapeutic outcomes. But, tumors are heterogeneous in their ability to repair damaged DNA, and evaluation population response only gives a statistical average. Thus selection of drugs without knowing tumor response at single cell level can cause side effects due to toxicity of drugs. There is a need to screen drugs reliably and rapidly for individual patients. This paper describes a new single cell based HaloChip assay that can be used to detect and quantify DNA damage and repair capacity after exposing to genotoxic drugs, and examine drug response without population interference. After forming cell array, cells are embedded in agarose that provides an interconnected network for DNA diffusion, followed by exposure to NaOH and stained with ethidium bromide. Dimensions of halo and nucleus are derived from collected fluorescent image. In the case of repair, cells are washed after exposing to drug and incubated for different time before HaloChip assay, the level of DNA damage is quantified using relative nuclear diffusion factor (rNDF) derived from surface areas of halo and nucleus. The rNDFs increase from 0 to 6 as drug concentration increase from 0 to 50 μM, and reach plateau when drug concentration is over 10 μM. At same dose, TOP-16 induces more DNA damage in HeLa cells than that of other two drugs; CPT-11 induces more DNA damage in LNCap cells than that of other two drugs. The bimodal repair curves are attributed to diversity of DNA lesions induced by drugs. The repair data are regressive fitted using first order exponential
New High-Throughput Version of the DEL Assay Detects Nonmutagenic Carcinogens.

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Genetic instability is a hallmark of carcinogenesis. Furthermore, cells from patients carrying mutations conferring cancer prone phenotypes show a higher level of genetic instability, including DNA deletions. In fact, the original yeast-based DEL (deletion) Assay with 100 chemicals shows an accuracy of 92% to detect carcinogens as compared to 62% detected with the Ames Assay. DEL events in all three formats are inducible by a wide variety of carcinogens including carcinogens that are negative in many other short-term tests. The DEL assay results also highly correlate with the clastogenicity of chemicals. Here we introduce the next generation DEL assay: a novel dual-read out Saccharomyces cerevisiae screen DEL-XG. DEL-XG simultaneously assesses the compound’s genotoxicity and cytotoxicity properties. During an exposure to a genotoxic event the sequence is excised and the lacZ gene recombines to a functional beta-galactosidase genotype that with an addition of the X-Gal substrate produces a strongly positive indole (blue) product. The most important advance is that we can determine the effect of target gene expression on genotoxicity. Surprisingly and most importantly we found that the Ames negative carcinogens that were weakly positive in our standard His reversion assay, are 10 fold more potent in expressed DNA. This means the new version of the DEL assay even more useful and makes sure that the Ames negative carcinogens will be positive even in the high throughput version of DEL-XG that is rapid and economical to screen large chemical libraries for toxicity and potential carcinogenicity properties.

In Vitro Genotoxicity Assays (Comet and Micronucleus) Using Engineered Skin.

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An in vitro micronucleus assay using human engineered skin and target cells grown beneath the tissues was developed. The purpose was to bring some information on exposure in vitro genotoxicity assays for dermally applied compounds. Previous results have shown that this method was reproducible and could be transferred to other laboratories. The system has now evolved to combine both the comet assay and the micronucleus assay. The approach is based on performing the comet assay in cells dissociated from the tissues, while the micronucleus assay was performed using the cells cultured beneath the reconstructed tissues. Four different time schedules were considered for this project: a 4 h treatment and a 27 h treatment period with or without a 27 h recovery period. A set of 13 chemicals were tested with this approach. The results obtained show that the best prediction model was the long treatment period (27 h) without recovery for both the comet assay and the micronucleus assay. Most of the ‘irrelevant positives’ yielded negative in vitro results using this system.

Development and Validation of a Toxicogenomics Signature to Differentiate Genotoxins vs Nongenotoxins.


Genotoxicity testing has long been used to assess a compound’s potential to induce genetic damage and hence it’s carcinogenic potential. The standard genotoxicity testing battery was designed to have high sensitivity but low specificity for detecting carcinogens that act by damaging DNA. As a consequence, positive genotoxicity test results require follow-up in vitro and in vivo testing to determine the biological relevance of the in vitro genotoxicity potential. Recent literature suggests that Toxicogenomic approaches could serve as a useful tool to identify different mechanisms through which compounds/ chemicals exerts their genotoxic effects. Although multiple research groups to date have published papers listing several gene expression based molecular signatures in vitro using human and rodent cell lines, and in vivo using rodent models, there exists no consensus largely due to species differences and diverse ‘omics’ technologies used. The broad objective of this study is to extend the work conducted by the ILSI-HESI technical committee on differentiating genotoxic and non-genotoxic carcinogens using real-time PCR (qPCR) based toxicogenomic approaches. In this study using a human lymphoblast TK6 cell line, we evaluated the utility of a qPCR technique to identify a molecular signature by studying gene expression profiles of twenty five genes. In our initial screening study we identified six genes that differentiated genotoxins vs non-genotoxins. However, in our validation study using ten model compounds per each group of aneugens, clastogens and non-genotoxins, we succeded in reducing six gene signature to three gene signature. The proposed three genes that could effectively differentiate clastogens and aneugens from non-genotoxins with high specificity and sensitivity are Cyclin-dependent kinase inhibitor 1A (CDKN1A), Growth differentiation factor 15 (GDF15) and Tumor protein p53 inducible protein 3 (TP53I3).

Characterization of Threshold Dose Response of Genotoxicity from Chemicals with Diverse Mechanisms of Damage.

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There is much debate regarding the existence of threshold dose-response for genotoxicity with DNA reactive chemicals. This study used in vitro high content imaging and flow cytometry assays together with Lutz threshold model (Lutz et al, 2009) to identify the threshold of dose-response for induction of double strand breaks (DSBs) and micronuclei (MN) in human p53-competent fibrosarcoma cells (HT1080). We evaluated 9 prototype chemicals: nocodazole (NCS; direct DSB inducer, mimics γ-irradiation), etoposide (ETP; topoisomerase II inhibitor), mitomycin C (MMC; DNA crosslinker), methyl methanesulfonate and ethyl nitrosourea (MMS, ENU; alkylating agents), hydrogen peroxide and tert-butylhydroquinone (H2O2, TBHQ; oxidative damage), quercetin and curcumin (QUE, CUR; oxidative polyphenols). All the oxidative agents H2O2, TBHQ, QUE and CUR induced a significant threshold response in both DSBs and MN, while the topo II inhibitor (ETP) and DNA crosslinker (MMC) induced non-threshold DSBs and MN response. The two alkylating agents showed different dose response trends: MMS exhibited threshold behavior of MN and DSBs, while ENU did not at the doses examined. Interestingly, NCS, a direct genotxin and potent inducer of DSBs, induced a linear-like DSB response, but a threshold increase in MN, indicating that the DNA repair response prevents conversion of DSB to MN at low doses. We also assessed p-p53 (ser15) and total p53 induction by these chemicals. The shapes of the p-p53 curves are generally comparable to p-H2AX, demonstrating that dose-response relationship of p53 activation follows similar pattern as DSBs. Our results indicate that dose-response relationship for DNA damage and MN formation may exhibit threshold or non-threshold depending on the chemical. Whole genome transcriptomics and repair enzyme proteins induction are currently being measured to evaluate the role of chemical specific activation/inhibition of DNA repair in the determination of linear vs. nonlinear dose-dependence of genotoxic response.


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Integrated testing strategies involve the assessment of multiple endpoints within a single toxicity study and represent an important approach for reducing animal use associated with testing. The study presents an evaluation to combine general, immune, and genetic toxicity endpoints into a single study. Specifically, this study evaluated the impact of sheep red blood cell (SRBC) immunization, as part of the T-cell dependent antibody response (TDAR) assay, on organ weights, RBC micronuclei formation, Comet response, and spleen T-cell immunophenotyping. Groups of female F344/DuCaP rats were dosed with cyclophosphamide (CP) by i.p. injection for five consecutive days at 0, 1, 3, and 10 mg/kg bw/day. Six rats from each dose group were injected i.e. with SRBCs on the initial day of dosing, as per the TDAR assay, while an additional 6 rats per group did not receive SRBCs. A
sham control group was included to account for animal handling and an additional group was dosed with ethyl malonate by oral gavage as a positive control for the comet assay. For the TDAR assay, treatment with CP resulted in a dose-dependent decrease in the antibody response with a suppression of greater than 95% at the high dose. Injection with SRBC had no impact on evaluated organ weights. Analysis of micronuclear formation revealed a dose-dependent increase in response to CP treatment, with an induction of greater than 5-fold at the mid and high doses. Injection with SRBC had no impact on the level of micronuclei in control animals and did not alter the dose response to CP. There was no increase in liver DNA damage in response to CP as measured by the comet assay and injection with SRBCs did not alter this endpoint. Similarly, injection with SRBC did not alter the spleen T-cell profile in response to CP. Overall these data provide strong support for the concurrent assessment of general, immune, and genetic toxicology endpoints within a single study as part of an integrated testing strategy approach.

2445 Evaluation of Repeated Dose Liver Micronucleus Assay in Rats: Summary of Collaborative Study by CSGMT/JEMS-MMS.


The repeated dose liver micronucleus (RLDMN) assay has a potential to detect genotoxic hepatocarcinogens that can be integrated into a general toxicological study. We have conducted a joint research in the Collaborative Study Group for the Micronucleus Test (CSGMT) to investigate the inter-laboratory variability and stable data acquisition in the RLDMN assay, which is supported by 19 Japanese facilities. In order to evaluate the performance of the assay, 28 chemicals including hepatocarcinogens were tested in 14- to 28-day RLDMN assays. As a result, the RLDMN assay detected the 9 chemicals positive out of 10 hepatocarcinogens, which were positive in the in vitro study while negative in the established bone-marrow micronucleus test. Also, the RLDMN assay detected the 2 hepatocarcinogens positive, which were positive in the short-term bone-marrow micronucleus test which was administered in the 14- and 28-day repeated dose liver bone marrow test. Accordingly, the RLDMN assay is not only useful in detecting genotoxic hepatocarcinogens but also appropriate for evaluation using a repeated low-dose regimen, and thus is considered ideal for integration into the general toxicity study.

2446 Acquisition of In Vitro Mutation and Cytogenetic Damage Information to Support Cancer Risk Assessment.

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Assessing the mode of action of carcinogenic chemicals is a critical component of cancer risk assessment. The US EPA’s Cancer Risk Assessment Guidelines stress the importance of determining whether or not a chemical causes tumors through a direct DNA-reactive mechanism, particularly in respect to the choice of an appropriate quantitative model for the extrapolation of cancer risk to low doses. Weight of evidence (WoE) approaches that rely heavily on data from in vitro hazard identification assays are commonly used to address carcinogenic mode of action. In vitro data needed to improve the WoE include both concordance analysis of temporal mutation induction and dose-response concordance of mutation with tumor incidence (e.g., Moore et al., Reg Toxicol Pharmacol 51:151-61, 2008). We used flow cytometric methods to acquire such data and thereby to test and extend this hypothesis. For example, male Sprague Dawley rats were treated for 28-consecutive days with the genotoxic carcinogen melphalan at 0.0, 0.031, 0.094, 0.28 and 0.75 mg/kg/day, which included dose levels corresponding to 0.3X, 1X, and 3X the tumorigenic dose rate of 50 (TD50). We assessed two endpoints of genotoxicity via low volume blood draws in order to efficiently obtain temporal data within the tumorigenic dosage range: MN assay at days 4 and 29, and Ploidy gene mutation at days 15, 29, and 56. The earliest time points evaluated showed dose-related increases for both endpoints, long before tumors or pre-neoplastic lesions would be expected. These data illustrate the potential of these blood-based analyses to provide dose-response and temporality information that relates genetic damage to cancer induction. Continuing studies will include additional genotoxic and non-genotoxic carcinogens and non-carcinogens to provide a deeper understanding of the relationships between genetic damage and cancer induction.

2447 Axonal Degeneration of Chronic Organophosphate Ester-Induced Delayed Neurotoxicity (OPIDN) Has Different Features in Central and Peripheral Levels of the Nervous System.

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OPIDN is considered one of the toxicant-induced central-peripheral distal axonopathies. The latter are characterized by degeneration of distal fibers in both the central and peripheral regions of the nervous system. This has been noted in a rat model of long-term exposure to tri-ortho-tolyl phosphate (TOTP). In the present report we draw attention to a difference in the nature of the axonopathy in the peripheral and central extensions of somatosensory fibers arising from neurons of dorsal root ganglia as seen in the sural nerve and spinal-medullary levels of the gracile fasciculus. As reported earlier (Jortner et al., Toxicol. Pathol. 33:378) young adult male Long-Evans rats were administered 14 TOTP gavage doses at 75, 150 or 300 mg/kg, over a 63-day period. Sacrifice was on days 63 and 90 (after a 27 day recovery period). OPIDN was manifest by TOTP dose-related diminished activity of brain neurotoxic esterase on day 63, and distal gracile fasciculus and peripheral nerve (including sural nerve) myelinated fiber degeneration. In the present report we draw attention to a qualitative difference between lesions of central somatosensory myelinated fibers in the distal levels of the gracile fasciculus and those in peripherally directed sural nerve fibers. Axonopathy progressing to myelinated fiber degeneration was seen in both regions, and was more florid in the gracile tract. In addition, prominent dystrophic axons (Jellinger, Prog. Neuropathol.,1973) were seen in the central region, and were absent peripherally. Axon dystrophy is considered to reflect terminal degeneration with retrograde progression, possibly due to aberrant regeneration, synaptic dysplasia, failing terminal catabolism or transport. This varying pathological response of myelinated fibers from the same neuronal population is considered an effect of the differing (central vs. peripheral nervous system) environments on the evolution of axonal lesions in this chronic neurotoxic condition. Supported by USAMRIC DAMD17-99-1-5489.

2448 Maternal Paraoxonase (PON1) Status Modulates Fetal Effects Associated with Gestational Exposure of Mice to Chlorpyrifos Oxon.

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Paraoxonase-1 (PON1) status (PON1 level and presence of the Q192R polymorphism) is an important determinant of toxicity for chlorpyrifos (CPF) and its metabolite, CPF-oxon (CPO). We examined whether maternal PON1 status influences fetotoxicity associated with gestational CPO exposure by comparing CPO toxicity among PON1-/-, wild-type (WT), and humanized transgenic mice expressing tgHuPON1R192 or tgHuPON1Q192. Pregnant mice were exposed dermally to 0.50, 0.75 or 0.85 mg/kg/d CPO from gestational days 6-17, and sacrificed on day 18 to measure enzyme inhibition in maternal and fetal tissues and gene expression in the GD18 fetal brain using Affymetrix microarrays. Fetal body weights from the PON1-/- dams exposed to 0.75 mg/kg/d CPO were significantly lower compared to vehicle controls. Pregnancy rate, number of resorptions and presence of fetal abnormalities were not significantly different among treatment groups in all genotypes. In the dams, repeated CPO exposure was associated with fetotoxicity. In the GD18 fetal brain, AChE expression was decreased, with the most apparent difference between PON1-/- and WT, while BChE expression was increased in these groups compared to WT.
Comparative Effects of Parathion and Chlorpyrifos on Extracellular Endocannabinoids: Influence on Cholinergic Toxicity.

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Parathion (PS) and chlorpyrifos (CPF) are organophosphorus insecticides (OPs). Acute subcutaneous exposure to either can elicit extensive acetylcholinesterase inhibition but cholinergic signs are relatively minimal following CPF compared to PS. Acetylcholinesterase (AChE) inhibition in vitro and moderate OP toxicity in vivo (Toxicol in Vitro, 25, 301, 2011; SOT abstract 2576, 2011: SOT abstract 942, 2012). Mechanisms contributing to the interaction of OP compounds such as paraoxon (PON) and fullerenes were evaluated. Formation of a covalent bond was ruled out by monitoring the release of p-nitrophenol and the decrease in concentration of PON by HPLC. Nuclear Magnetic Resonance (NMR) was used because hydroxylated fullerenes are known to make aggregates in aqueous media and PON could be complexed with two hydroxylated fullerenes. Using reversed phase high performance liquid chromatography (RP-HPLC) and Nuclear Magnetic Resonance (NMR), we have identified PON by HPLC. Nuclear Magnetic Resonance (NMR) was used because hydroxylated fullerenes are known to make aggregates in aqueous media and PON could be complexed with two hydroxylated fullerenes.
after 15 min incubation with the C80 derivative containing gadolinium (GdTMSOH) and a C70 derivative (C70-OH) without any metal. Proton-decoupled phosphorus (31P) NMR spectra were recorded on a Bruker Advance III 600 NMR instrument. The experiments with PON and the fullerene were run in deuterium oxide with phosphoric acid 0.04% as an internal reference. Results for GdTMSOH demonstrated a small upfield change in the chemical shift as the concentration of fullerene increased. However, due to strong interferences from the metal, further studies were done with the C70-OH fullerene. Plotting the variation of its chemical shift versus the concentration of fullerene resulted in a binding isotherm curve that reached a plateau at 0.03 mol/l. The binding constant (Kb) calculated by dividing the intercept by the slope from the double reciprocal plot was determined to be 19 (mol/l)-1. The results suggest that the fullerenes act in a manner much like cyclodextrins, which have previously been shown to chemically interact with OP compounds, decreasing their capability to inhibit AChE (Carbohydrate Res. 345, 141, 2010; Euro J Med Chem. 40, 615, 2005; Toxicology 265, 96, 2010). Supported by the CounterACT Program, NIH OD and NINDS, grant NS063723.

### 2454 Conditional Toxicity Value (CTV) Predictor for Generating Toxicity Values for Data Sparse Chemicals.

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Chemical hazard assessments necessarily vary based on data availability and the type of risk management decision they support. While much recent attention has been on use of high-throughput toxicological data for screening and prioritization, assessments that support toxicity guidance values or standards still rely on epidemiological and in vivo experimental data. Such assessments, including Integrated Science Assessments and Integrated Risk Information System Toxicological Reviews, are highly data-, time-, and resource-intensive, and cannot be realistically expected for most environmental chemicals. Thus various stakeholders and expert groups, including the National Research Council in Science and Decisions, call for "default approaches to support risk estimation for chemicals lacking chemical-specific information." This project aims to address this challenge through the Conditional Toxicity Value (CTV) Predictor. This tool uses chemical properties and limited experimental data to predict toxicity values, such as the reference dose (RfD) and concentration (RfC), oral slope factor (OSF), inhalation unit risk (IUR), or cancer potency value (CPV). CTV predictions combine QSAR, regression, and hybrid modeling, rely on a new comprehensive database of existing guidance values and experimental data, and incorporate OECD principles for model building and external cross-validation. QSAR models for predicting existing RfD values (which span 7 orders of magnitude) had an R2 up to 0.46 and Absolute Error of 0.67 ± 0.03 (Log10 mg/kg/day). A tool that can predict a toxicity value within an order of magnitude fills a critical gap in the current risk assessment methodology.

Disclaimer: The views expressed here are the authors' and not necessarily those of the US or California EPA.

### 2455 Application of the Threshold of Toxicological Concern (TTC) Decision Support Approach to Antimicrobial Pesticides.

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A tiered decision support approach has been developed to apply existing knowledge of antimicrobial pesticides (AMPs) to new AMPs being considered for use. The approach is intended to inform product development and regulatory review processes so that antimicrobial pesticides and pesticide products can be targeted early in development and so that animal testing can be focused on those chemicals that need the most attention. Expert groups were convened comprised of scientists from non-government organizations, industry, academia, and government. The experts have 1) collected high quality data from studies submitted to EPA’s Office of Chemical Safety and Pollution Prevention and the U.S. Food and Drug Administration’s regulatory review files and entered the data into a publicly accessible data set using the structure and approach of EPA’s ToxRef Database; 2) developed a tiered decision framework for estimating systemic dose from dermal exposure so that comparisons could be made to the oral toxicity data available; 3) applied chemoinformatics techniques to evaluate whether and how to bridge the AMP data set to the 3 non-cancer TTC values of Munro et al (1996); 4) also used chemoinformatics techniques to define classes of AMPs within a TTC decision framework, and 5) developed a decision tree to guide consideration of new AMPs with respect to likely toxicity for anticipated systemic dose of a given formulation and the chemoinformatics class into which it falls. We will present the method for data curation, chemoinformatics, rationale for the decision approach, and case examples of the use of the approach. This work will advance the science and practical use of computational toxicology in support of risk management decision making.

### 2456 The Navigation Guide As an Evidence-Based Medicine Methodology to Evaluate Human Health Effects of Environmental Chemicals: Perfluorooctanoic Acid (PFOA) and Fetal Growth.

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Rationale: Evaluating environmental health literature and determining the weight of evidence are critical for informing policy and health recommendations. The National Academy of Sciences has called for an enhanced systematic and transparent approach to risk assessment and scientific decision-making. The Navigation Guide was developed through a collaboration of 22 scientists to improve methods of research synthesis in environmental health. The methodology is based on best practices in evidence-based medicine and environmental health sciences and aims to systematically and transparently synthesize the evidence from toxicology and observational epidemiology studies.

Approach: To establish proof of concept, we applied the Navigation Guide to the question of the impact of exposure to perfluorooctanoic acid (PFOA) on fetal growth. Steps include: (1) Specify the study question; (2) Select the evidence; (3) Rate the quality and strength of the evidence.

Findings: We identified 24 human observational and 21 animal toxicological studies relevant to the study question. Study quality was assessed using a modified version of the Cochrane Collaboration's Risk of Bias tool. Preliminary meta-analysis of combinatorial studies suggests there may be small reductions in birth weight with increased PFOA exposure in animals and humans.

Implication: The case study illustrates that the Navigation Guide can be used to apply the rigor of systematic review methodology to questions in environmental health. As has been demonstrated in the clinical field, the adoption of a systematic and transparent method to synthesize the scientific evidence in the environmental health field would speed incorporation of research into decision-making.

### 2457 Incorporating Population Variability and Susceptible Subpopulations into Dosimetry for High-Throughput Toxicity Testing.

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 Xenobiotic clearance can vary widely across age-based or ethnic subpopulations due to differences in metabolic enzyme abundances and activities. A strategy to measure population-specific hepatic clearance values across a wide range of chemicals would allow pharmacokinetic variability due to age, ethnicity, and other factors to be incorported into in vitro toxicity screening data. Metabolic clearance of ToxCast chemicals selected based on LC-MS method compatibility and exposure estimate availability were measured in vitro using 13 cytochrome P450 (CYP) and 5 UDP-glucuronosyltransferase (UGT) recombinantly expressed isoforms. Together with plasma protein binding, these isoform-specific clearance rates were then incorporated into an in vitro-to-in vivo extrapolation (IVIVE) modeling tool, Simcyp. The modeling tool accounts for known differences in isoform abundances among various age- or ethnic-based subpopulations to estimate the daily oral dose for each subpopulation, called the oral equivalent dose, necessary to produce steady-state in vivo blood concentrations equivalent to in vitro AC50 values across the -600
2458 Extrapolating In Vitro Embryotoxicity Data Toward In Vivo Exposure Levels Using a Combined In Vitro-Physiologically-Based Kinetic Modeling Approach.

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In vitro assays play an important role in screening chemicals for their toxic potency and prioritizing them for further toxicity testing. Most of these in vitro assays are not suitable for a quantitative risk characterization as they lack in vivo kinetic processes. To overcome this limitation, the present research aimed at combining in vitro toxicity data with physiological based kinetic (PBK) modeling to predict in vivo exposure levels. In order to contribute to the 3Rs principle for the replacement, reduction and refinement of animal testing in the most optimal way the required PBK models were developed on the basis of only in vitro and in silico data and data available from literature. Phenol was selected as the model compound and the end-point of interest concerns embryotoxicity. At first, the embryotoxicity of phenol was evaluated in vitro using the embryonic stem cell test (EST), revealing a concentration dependent inhibition of differentiation into beating cardiomyocytes. In a second step, PBK models were developed for rat and human using in vitro derived kinetic constants, in silico derived physico-chemical parameters and literature derived physiological data. After evaluating the performance of the PBK models, the in vitro concentration-response information from the EST served as an input in the PBK model, thereby creating an in vivo dose-response curve from which a Point of Departure for risk assessment could be derived. Finally, in vitro embryotoxicity effect levels available from literature were used to evaluate this combined in vitro-PBK approach. In summary, this study shows how combining different alternatives in vitro toxicity testing and PBK modeling will enlarge their application from screening and prioritizing of chemicals toward deriving safe exposure levels for the risk assessment of chemicals.

2459 A Genomics-Based Determination of Relative Potencies of Dioxin-Like Compounds in Primary Human Hepatocytes.

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Toxic equivalency factors (TEFs) for dioxin-like compounds are predominantly derived from relative potency (REP) determinations of endpoints such as enzyme activity from animal studies and in vitro studies in immortalized animal cells. Currently, REPs based on gene expression changes have not been considered when deriving TEF values. Moreover, data from humans and human cells are not included in the REP database for deriving TEF values. In this study, primary human hepatocytes were treated for 24 hours with 11 concentrations of 2,3,7,8-tetrachlorodibenzo-p-dioxin, 2,3,4,7,8-pentachlorodibenzofuran, or 2,3,7,8-tetrachlorodibenzofuran ranging from 0.00001-100 nM. Gene expression changes were analyzed using ANOVA to assess the relative contributions of concentration, congener, and the interaction between concentration and congener. A total of 1,445 genes showed significant changes with concentration (FDR < 0.05 and fold-change +1.5 in at least one concentration for one congener). Of these, 399 were significant for both concentration and congener effects indicating parallel concentration response curves with differences in potency. Pathway enrichment analysis was performed on the 1,443 genes differentially expressed by concentration. The top 10 most enriched pathways included several nuclear receptors, immune response, and cell adhesion pathways. A focused assessment of the benchmark dose values for signaling pathways in the modes of action for DLC-induced effects can be used to determine more relevant REPs used by the World Health Organization (WHO) to determine TEF values. The addition of human cells to the WHO database could provide more insightful measures of relative potency and provide quantitative data to reduce TEF uncertainty in human health risk assessments.


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XDE-729 methyl, a novel herbicide in development, induces rodent liver enlargement, hyper trophy, and hyperplasia via an aryl hydrocarbon receptor (AhR) mediated mode-of-action (MoA) with the following key events: 1) pre-systemic liver exposure to XDE-729 methyl, 2) AhR activation with associated liver hypertrophy, leading to 3) hepatocellular proliferation. For key event 1, low levels of XDE-729 methyl are present in the liver after dietary exposure; however, XDE-729 methyl is rapidly metabolized in the liver to primarily XDE-729 acid, which is not an AhR activator, and excised in the urine. For key event 2, AhR activation, measured by Cyp1a1 transcript induction occurred at dose levels of ≥50 mg/kg/day XDE-729 methyl and correlated with liver hypertrophy. For key event 3, hepatocellular proliferation was observed with exposure to 261 mg/kg/day XDE-729 methyl. A threshold for AhR activation and liver effects occurs at 10 mg/kg/day XDE-729 methyl, the no-observed-adverse-effect level (NOAEL) from the rat 90-day toxicity study. Collectively, the data provides a high level of confidence that XDE-729 methyl induces through the rat liver, but not an AhR dependent MoA at doses ≥10 mg/kg/day XDE-729 methyl. The AhR pathway is conserved across species and prototypical AhR ligands are activators of both human and rodent AhR; however, human AhR binding affinity for prototypical AhR ligands is quantitatively lower than rodent AhR. XDE-729 methyl is rapidly metabolized to XDE-729 acid, which does not activate AhR, and does not bioaccumulate in the rat liver. XDE-729 methyl does not result in sustained activation of the AhR pathway and hepatic effects are transient and reversible. Based on the above, a margin of exposure risk assessment using the 10 mg/kg/day NOAEL from the rat 90-day toxicity study to derive the chronic reference dose/acceptable daily intake (cr/ADI) for XDE-729 methyl is protective of human health.

2461 Estimation of Cumulative Risk from Exposure to Phthalates Using NHANES Exposure Data and an In Vitro Potency Assay.

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Endocrine active phthalates reduce testosterone synthesis in fetal rat testes, interfering with male sexual development at high doses. As they share the same mode of action, it is logical that a phthalate risk assessment should account for cumulative exposure. Lack of potency data in vivo limits the derivation of a cumulative risk estimate. We illustrate an approach using in vitro pharmacodynamic data together with NHANES exposure estimates to predict human risk from multiple phthalates. First, we developed a rat Leydig tumor cell line (R2C) in vitro assay to assess testosterone inhibition with several environmentally relevant phthalate metabolites: monobutyl (MBP), monooctyl (MOEH), monoethyl (MEP), monomethyl (MMP), monononyl (MOP), monobenzyl phthalate (MBzP) and two oxidative metabolites of MEHP (5-Ox-MEHP, 5-OH-MEHP). Effects of these chemicals on steroidogenesis in vitro were highly consistent with in vivo studies. In vitro IC50 values were used to derive relative potency factors (RPFs) in relation to MBP. These RPFs were used with the 2005 NHANES urine data to calculate relative risk estimates for each phthalate at the 50th or 95th percentile (exposure). Estimates for individual phthalates were then combined to obtain a dibutyl phthalate (DBP) equivalent risk estimate (mg DBP/kg/day). This in vitro predicted cumulative risk estimate was then compared to the USEPA in vivo derived reference dose (RfD) for DBP. At the 50th percentile DBP accounted for 8% of total exposure but represented 70% of the total risk of all phthalates. At the 95% however, the other phthalates contributed more to the cumulative risk. Cumulative potency weighted exposure for all phthalates was markedly lower than the USEPA RfD for DBP (0.1 mg/kg BW/day), whether calculated for exposure at the 50th (0.8007 mg/kg/day) or 95th (2.7860 mg/kg/day) percentile. Exposure for all phthalates was markedly lower than the USEPA RfD for DBP (0.1 mg/kg BW/day), whether calculated for exposure at the 50th (0.8007 mg/kg/day) or 95th (2.7860 mg/kg/day) percentile.
2462 Nonhuman Primate Sexual Maturity: What Is the Capacity to Endure Uncertainty?
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Evaluation of reproductive toxicity in regulatory studies relies on the use of sexually mature animals. This often involves the use of nonhuman primates (NHP) when lower order species are pharmaceutically irrelevant. However, the decision on whether to utilize mature NHP and the criteria used to establish sexual maturity is anything but standardized. Several factors contribute to these differing opinions. Sexual maturity can occur over a wide range of ages and body weights and historical methods of predicting sexual maturity have not always correlated with the histopathological appearance of the gonads at necropsy. Evaluation of toxicity in the reproductive organs of pre/peripuberal animals is often difficult, given that the histology of the maturing reproductive organs may resemble the degenerative changes induced by reproductive toxicants in the sexually mature adult. Finally, there is a lack of consensus on the toxicological relevance and interpretation of the various possible reproductive endpoints. All of these factors result in a complicated balancing act: Ethically weighing the limited availability, high costs, and relevance of incorporating sexually mature NHP in toxicity studies against effective and meaningful evaluation of reproductive risks. This roundtable will discuss the challenges of assessing toxicity in immature vs. peripuberal vs. mature NHP, screening methods for identifying sexually mature NHP, factors that influence sexual maturity, and case studies where sexual maturity impacted the study interpretation. Overall, this session seeks to provide an opportunity for stakeholders to review the current state of the art and exchange views on appropriate paths forward to encourage ethical use of animals, while preserving appropriate risk assessments.

2463 Exposure Science in the 21st Century: Perspectives from the NAS and What It Means for Toxicology.
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In 2010, US EPA with additional support from NIEHS, requested the National Academy of Sciences (NAS) to develop a long-range vision for exposure science in the 21st century and a strategy for implementing this vision over the next twenty years. Exposure science is the bridge between the sources of chemical, physical and biological agents and ecological and human health. Exposure science is critical for predicting, preventing, and reducing human health and ecosystem risks. The report, Exposure Science in the 21st Century: A Vision and a Strategy, was released September 7 by the NAS National Research Council. This report along with three other NAS reports, Toxicity Testing in the 21st Century, Science and Decisions: Advancing Risk Assessment and Sustainability and the US EPA, chart the future directions for using innovative technology and scientific advances to better understand environmental impacts on human and ecological health. The report outlines a framework for advancing exposure science to study how humans and ecosystems interact with chemical, biological, and physical agents in their environments. Key visions in the report in are to develop a universal exposure-tracking framework with a focus on preventing and mitigating adverse exposures. This will include the application of systems science to understand and characterize exposures across time, space, and biological scale.

2464 Regulatory-Based Nanotoxicology: Evolving National Strategies, and Research to Address Engineered Nanomaterial Health Risk Assessments.

Engineered nanomaterials are increasingly being developed and incorporated into a variety of products and applications. However, the full development of nanotechnology is hampered by an uncertainty regarding their environmental, health and safety implications, and how these issues will be addressed by responsible regulatory agencies at the federal and state levels. Novel nanoscale materials present a number of scientific and technical challenges for assessing their potential health implications including: exposure and material characterization; adequacy of conventional toxicity testing methods/guidelines; dose metric(s) across the exposure-dose-effects paradigm; and the role of alternative testing approaches to assess their toxicity. This session will bring together key stakeholders from scientific advisory bodies, regulatory agencies and the private sector to present/discuss research needs, strategies, approaches and findings regarding the health effects testing of engineered nanomaterials and nan-enabled products as it relates to regulatory mission(s). Key topics include: assessing nanomaterial toxicity for regulatory and risk assessment applications; the status or role of alternative test methods to screen or prioritize nanomaterials for in vivo toxicity testing, and the extent to which harmonized testing can be achieved for these novel materials. Each speaker will have 10-12 minutes to highlight agency/institutional approaches and regulatory actions regarding the potential for health effects from engineered nanomaterials. The session will conclude with a 15 minute general discussion period. The overall goal is to provide participants with a view of the status of the development of nanomaterial toxicological assessments that would be sufficient to evaluate health and safety for regulatory agencies.

2465 Toxicological Writing for Industrial and Regulatory Audiences.
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Excellence in scientific and technical writing leading to high-quality publications, a skill set developed and refined from graduate training through early career in toxicology, is one key trait that can lead to a successful career as a toxicologist. Some academic institutions have programs that support scientific and technical writing for their faculty. However, the majority of toxicologists (80%) are employed outside academia, predominately within biopharmaceutical and chemical industries, government, and contract research organizations. Graduates from academic programs that train in writing might acquire skills related to preparation of dissertations, grant proposals, and manuscripts for scientific peer-reviewed journals. The skills acquired from this training, when existent, does not necessitate a smooth transition to a successful career outside academia. Thus early-career toxicologists are sometimes unaware of, or otherwise unprepared for, technical writing assignments that occur in industrial and regulatory toxicology. In addition, different writing skills are required for clear and concise communication of toxicological results to nontoxicologist stakeholders. Therefore, an interactive workshop that evaluates the challenges presented by, and the skills required for, toxicological writing outside academia will be of great use to the majority of graduate students, postdoctoral trainees, and early-career toxicologists. Four speakers from the pharmaceutical industry, the chemical industry, a government regulatory agency, and a contract research organization will review the type(s) of technical writing required within their setting. Notably, they will extensively highlight common mistakes and discuss valuable strategies and tools to avoid these mistakes through interactive exercises using provided writing examples.

N. J. Walker, National Toxicology Program, NIEHS/NIH, Research Triangle Park, NC.

In recent years there has been considerable research examining the potential utility of nanoscale materials and nanostructures in commercial and biomedical applications. Nanoscale materials (nanomaterials, nanoparticles), are a broadly defined set of substances that have at least one critical dimension less than 100 nanometers. The same novel chemical and physical properties that make nanomaterials useful also make their interactions with biological systems difficult to predict and evaluate in traditional toxicity models. The diversity in composition, size, surface coatings, and physico-chemical properties even within classes of “similar” nanomaterials can make translation of findings from one nanomaterial to another problematic. On the other hand though, testing of each nanomaterial individually and what change in a physico-chemical property constitutes creation of a “new” material is uncertain. Over the past years there has been considerable effort looking at developing guidance for assessing safety of nanomaterials. Some of the key considerations include: effective characterization of physicochemical properties of nanomaterials not only in the bulk phase but also within the toxicological test system; use of in vitro biological responses and shorter term in vivo studies of panels of nanomaterials and nanostructures in commercial and biomedical applications. In recent years there has been considerable research examining the potential utility of engineered nanomaterials (ENMs) has increased significantly in recent years due to potentially hazardous impacts on human health. Mast cells are critical for innate and adaptive immune responses, often modulating allergen-induced pathologic conditions. Mast cells are well known to act in response to danger signals through a variety of receptors and pathways including IL-33 and the IL-1 like receptor ST2. We have examined the involvement of mast cells and the IL-33/ST2 axis in pulmonary and cardiovascular responses to ENMs including multi-walled carbon nanotubes (MWNT) and silver nanoparticles (AgNP). We have utilized C57BL/6 mice, mast cell deficient mice (KitW-sh), KitW-sh mice reconstituted with wild-type or ST2–/– mast cells and ST2–/– mice to assess systemic and pulmonary inflammatory responses as well as cardiac ischemia-reperfusion (IR) injury responses following ENM exposure. In addition, we have used an in vitro mast cell model to screen for the ability of ENMs to directly induce mast cell degranulation. We have found that mice with normal mast cell populations (C57BL/6 and mast cell reconstituted KitW-sh) exhibit significant ENM directed systemic and pulmonary inflammation, fibrosis, altered lung function and exacerbated IR injury. In contrast, these toxicological effects of ENMs were not observed in mice deficient in mast cells (KitW-sh) or mice with mast cells unable to respond to IL-33 (ST2–/– mast cell reconstituted KitW-sh mice). Lastly, we have established that certain ENMs are capable of inducing mast cell activation in vitro. Our findings establish for the first time that mast cells and the IL-33/ST2 axis orchestrate adverse immune effects to ENMs giving insight into a previously unknown mechanism of toxicity and providing a realistic therapeutic target. The use of mast cells and the IL-33/ST2 axis as a screening tool for ENM toxicity and in the preclinical development of nanomedicines will be discussed.

Molecular Dynamics Simulations with Advanced Sampling Techniques to Study Nanoparticle-Membrane Interactions.

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Classical toxicology assessments consider bulk transport of particulate matter into discrete organelles of living cells at the primary means of nanoparticle (NP) toxicity. However, little is known about the potential of low-level exposure to alter physiologic functions (membrane form/function). Atomistic simulations, such as molecular dynamics (MD), can provide mechanistic information that is hard to measure experimentally. They may be used to validate theories derived from indirect experimental analysis, but the results of MD simulations are meaningful only if the run is long enough to visit all energetically relevant configurations. Well-tempered metadynamics can accelerate system dynamics allowing the analysis of processes that can take several seconds or hours to occur in real systems. Classically biased MD simulations were used to reconstruct the free energy landscapes of the mechanisms of NP entry into biological cells, thus accelerating the sampling of rare events and allowing exploration of potentially new biologically relevant reaction pathways. Results show that pristine C60 resides preferentially in the aliphatic region of the lipid bilayers composed of POPC and cholesterol. With increasing cholesterol concentration, the minimum of the free energy profile moves towards the interface with water, showing a tendency of C60 to move away from the membrane center. C60 mobility inside the hydrophobic region is not limited by any relevant thermodynamic barriers at body temperature, but the energetic cost to leave the membrane is system-dependent and varies from 10 to 20 kJ·mol. Charges (C60+) or hydroxyl groups (C60(OH)3) make the region where the lipid heads are solvated by water the most favorable position for these species. These studies make plausible the potential for intramembranous mechanisms of NP toxicity in immune and other mammalian cells. Supported by ES08846, NIEHS NCNHR U01 (MAP), CBET 0644639 (AV), 1 U02 ES020128-01 (KR), T32-ES007062-26 (KR).

Evaluating the Local and Systemic Immunomodulatory Effects of Nanomaterials.

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The small size of nanomaterials (NM) makes them a prime target for interaction with the immune system following their uptake, processing, and presentation to lymphocytes by antigen-presenting cells. Traditional in vivo testing strategies have been used to evaluate NM-mediated immunotoxicity. The varied routes of exposure to NM (dermal, oral, inhalation, parenteral), as well as particle size, can produce different immunological responses including contact sensitivity as well as local or systemic effects. For example, a single phylogeneric aspiration of anatase TiO2 nanoparticles (< 25 nm) produced an enhanced antibody-forming cell response to sheep erythrocytes, while TiO2 microparticles (< 45 μm) produced no such effect. Subcutaneous non-dermal, exogenous exposure to anatase nano-TiO2 for 3 days increased cell proliferation in the draining lymph nodes, while oral exposure for 28 days was non-immunotoxic. Inhalation of 1.0 and 0.05 μm C60 fullerene for 13 weeks did not affect the systemic immune response, although immunological cytokines (MCP-1, MIP-1α) in the bronchoalveolar lavage fluid were increased for the 1.0 μm C60 only. Successful implantation of a recombinant electroporation biodegradable poly-caprolactone materials in the ventral quadrants for 28 days had minimal effects on the immune system, suggesting that this material may have the potential for use in a variety of clinical applications. Nanomedicines intended for clinical applications unrelated to the immune system must be examined for the possibility that they might produce unintentional or unanticipated immune effects. However, depending upon the application (e.g. local anti-inflammatory drug delivery), effects on the immune system may be desirable.

Phenotypically Profiling the Factors Affecting the Pharmacokinetics and Pharmacodynamics of Nanoparticle Agents in Preclinical Models and in Patients.

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Carrier-mediated agents consist of nanoparticles, nanosomes (nanoparticle sized liposomes) and conjugates. The theoretical advantages for using carrier-mediated drugs, including increased drug solubility, prolonged duration of exposure, selective delivery of entrapped drug to the site of action, and improved therapeutic index. Pegylated nanosomal formulations contain lipid conjugated to polyethylene glycol. The disposition of encapsulated drug is dictated by the composition of the carrier, thus altering the pharmacokinetic (PK) profile of the drug. A proposed clearance pathway of carrier agents is the monocytes and macrophages of the mononuclear phagocyte system (MPS). Our studies suggest there is a bi-directional interaction between nanosomal agents and the MPS. However, potential factors associated with clearance of carrier agents in patients and preclinical animal models have not been extensively evaluated. Standard allometric scaling approaches for nanosomal agents did not scale across all species. In addition, preliminary studies suggest that there is high variability in the function of the MPS in animal models and in patients. Thus, new methods for allometric scaling and measures of MPS function need to be developed for carrier-mediated agents. In addition, the most appropriate animal models for toxicology and pharmacology of carrier agents needs to be identified. Novel methods for phenotypically profiling the factors affecting the pharmacokinetics and pharmacodynamics of nanoparticle agents in preclinical models and in patients will be presented.
Within a population of genetically heterogeneous individuals, a range of responses is observed for environmental exposures. The observed variability in response is attributable to extrinsic and intrinsic factors, including individual differences in exposure to environmental stressors and genetic/epigenetic heterogeneity, respectively. Current risk assessment practice is to account for interindividual variability with default uncertainty factors (e.g., a ten-fold decrease in allowable exposure to protect the most sensitive subpopulations), even though these defaults are seldom supported by scientific evidence. Advances in exposure science and molecular genetics are greatly increasing our ability to characterize intrinsic differences among individuals in their exposure and response to toxicants. This symposium highlights several novel and exciting approaches in safety evaluation that utilize recent advances in genetics. First, recent collaborative efforts in the complex traits community have led to the development of several new, powerful mouse resources that greatly facilitate the identification of allelic variants of genes associated with differential response to toxic exposure through genotype-phenotype associations. Second, several laboratories, including the National Toxicology Program, have begun applying these new mouse models of human population diversity to studies on the molecular mechanisms of interindividual variability in chemical metabolism and toxicity. Third, the ability to generate induced pluripotent stem (iPS) cells from population-derived human cell resources, as well as the availability of embryonic stem (ES) and iPS cells from mouse strains, makes it possible to conduct in vitro studies to investigate interindividual differences in resistance and susceptibility to xenobiotic exposures. Ultimately, these new approaches should greatly enhance our ability to characterize variability in response to toxicants and to identify those genes and pathways that contribute significantly to the observed differential responses to environmental exposures in humans.

Benzene is hematotoxic, genotoxic, and tumorigenic to the lymphohematopoietic systems in both laboratory animals and humans. Results from NTP population-based mouse models of benzene are consistent across studies, showing significant individual variability in benzene ADME/TK and levels of DNA damage at human relevant exposures. The results of these studies illustrate how individual variability in genetically defined populations may be used to identify genetic variants associated with differences in toxicokinetics, resistance or susceptibility to a toxicant, and to identify causally related mechanisms of toxicity. Benchmark dose models can be used on quantitative data to determine a reference dose that will aid in the quantification of uncertainty factors and, possibly, eliminate default assumptions in risk assessment. In additions, by identifying multiple genetic variants through genotype-phenotype associations with significant site effects, networks and pathways may be predicted that are statistically anchored to toxicity phenotypes and functionally-validated through the use of recombinant inbred mice (Collaborative Cross) cell-based assays, in vivo targeted testing, and molecular biology using reverse genetics. Across species extrapolation is enhanced by identifying variants that are orthologous between mouse and human and if there are associated in orthologous networks or pathways. In combination with high-throughput and high content assays validated by in vivo targeted testing, these new tools will provide a new paradigm for toxicology and exposure related diseases.

A genetically diverse panel of ES cell lines have been produced from inbred, F1, and outbred mice, and this panel has been used to investigate the role of strain background on variable response to environmental toxicants. An example of such a study is an investigation of how genetic background and environmental factors interact in the onset or severity of arrhythmia, and how these interactions may be involved in the cardiotoxicity of environmental toxicants. The ultimate goal is to uncover novel mechanisms, targets, and chemical structures underlying variability in cardiotoxic response. The approach is to conduct large scale, high throughput in vitro screens using cardiomyocytes derived from genetically diverse ES or iPS cells to interrogate gene toxicant interactions in vitro. Toward this end, a large panel of genetically diverse mouse ES lines as well as an automated beating assay performed in 384 well plate format for time domain analysis of arrhythmia have been developed. Data based on testing reference compounds of known arrhythmogenic potential in ES derived cardiomyocytes support the validity of this approach.

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the context of Parkinson’s disease. In addition, recent epidemiological findings link- ing residential proximity to freeway with autism will be presented. The intent of the symposium is to stimulate discussion especially in the context of future research needs in this rapidly emerging area of neurotoxicology which requires a multidisci- plinary approach involving expertise in the toxicology of the CNS, air pollution, mixtures, and nanoparticles in order to conduct appropriate risk assessment.

2479 CNS Consequences of Postnatal Exposure to UFP
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Accumulating evidence suggests that air pollutants can induce inflammation and oxidative stress in the brain and that early development may be a period of particu- lar vulnerability to such exposures. Epidemiological studies report associations be- tween air pollutant exposure and impaired cognitive function in children and adults (Tallarida and Block, 2002). Animal studies seek to determine the effects of developmental and/or adult concentrated ambient particulate matter (CAPS) expo- sure on cognitive behaviors and their potential CNS mechanisms. C57Bl6 mice were exposed to AIR or CAPS during early postnatal life with or without adult challenge, yielding 4 groups: AIR/AIR, AIR/CAPS, AIR/CAPS/AIR and CAPS/CAPS. Behavioral measures included repeated learning, a fixed ratio wait for reward para- digm related to impulsivity (males only). Fixed Interval schedule-controlled behav- ior and locomotor behavior. Learning accuracy was significantly impaired after CAPS/Air and CAPS/CAPS exposures, particularly in females, even though ability to perform already-learned responses remained intact. In the fixed ratio wait for re- ward paradigm, CAPS/AIR and CAPS/CAPS decreased the time animals would wait for free reward deliveries, and thereby resulted in emission of a greater number of responses for reach reward earned. Response rates on the fixed interval schedule were reduced in CAPS/AIR males but increased in Air/CAPS females. Notably, as measured at the termination of behavioral testing, i.e., almost a full year after the adult exposures, histopathological hallmarks of CNS perturbation, namely ax- ogliosis and increased microglial presence, were seen in multiple brain regions, in- cluding dentate gyrus and ventral midbrain and increased glutamate and GABA was prevalent in frontal cortex of both sexes. Current efforts focus on understand- ing the relationships between inflammatory and neurotransmitter changes and be- havioral outcomes. Exposure to CAPS appears to have long-term consequences for the CNS. As such, air pollutants may represent an underappreciated contribution to CNS disease and disorders.

2480 Microglia and the Peripheral Immune Response in Diesel Exhaust-Induced Neuropathology.
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Air pollution has been linked to central nervous system (CNS) disease, but the mechanisms driving this response and the type of exposures responsible are largely unknown. To discern the early CNS response to diesel exhaust (DE), adult male rats were exposed to DE (2.0, 0.5, and 0 mg PM/m3) for one month by inhalation. DE exposure elevated markers of neuroinflammation (nitrotyrosine, IL-6, TNFα, and MIP-1α) in the midbrain, cortex, and olfactory bulbs, in addition to activation of microglia, as determined by morphology. The highest levels of neuroinflamma- tion occurred in the midbrain, which also contained the highest level of the IBA-1 microglial marker. Neurotoxicity and early markers of neurodegenerative disease were unaffected. One month exposure to biodiesel exhaust revealed only changes in microglia morphology without any elevation of pro-inflammatory factors, demon- strating that chronic inflammation may be exposure specific. To reveal if DE might be causing neuroinflammation, rats were administered an IT bolus of DE particles (DEP 20 mg/kg) where DEP exposure elevated serum and brain TNFα, suggesting a potential role for the particles in initiating peripheral inflammation capable of transferring to the brain. Microglia cultures treated with DEP (90µg/ml) failed to produce TNFα, indicating that brain TNFα production may not be due to the di- rect interaction of microglia with the particles. To explore the role of long-term ex- posure, we tested the effect of subchronic (6 month) DE (992, 311, 100, 35 and 0 µg PM/m3) inhalation in rats and found elevated levels of the neurotoxic cytokine TNFα at lower exposures (DE, 100 µg PM/m3) and early protein markers of neu- rodegenerative diseases (α-synuclein and phosphorylated tau) at higher lev- els (DE 992 µg PM/m3), suggesting that subchronic DE exposure may impinge on common neurodegenerative disease pathways. Together, these findings demonstrate that the brain's immune system can detect and respond to DE and support that cir- culating cytokines (or another peripheral signal) may play a role in how neuroin- flammation transfers to the brain.

2481 Particle Translocation As an Explanation for the Adverse Effects of Inhaled Particulates in the CNS.
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The hypothesis that air pollution can adversely impact the central nervous system (CNS) is supported by the demonstration of gene-environment interactions for neurodegenerative disorders, such as Alzheimer’s and Parkinson’s diseases. The mechanisms by which this might occur are not fully understood, but may include transport of particulates into the brain, thus resulting in direct delivery of toxicants to the target tissue. Transport via the olfactory route, for example, has been demon- strated for polycyclic ultralite or nanosized particles (<100 nm in diameter in at least one dimension) and for some soluble compounds following inhalation expo- sures. However, other pathways have been identified and the contribution of these to the overall accumulation in the CNS is unclear. Likewise, the contributions of particle physicochemistry and blood-brain barrier integrity are important to ad- dress in order to understand the long-term consequences to CNS health. Such con- sequences have been explored in recent research. Using a mouse model of Alzheimer’s disease, we found that a 12-day inhalation exposure to poorly soluble manganese oxide nanoparticle aerosols resulted in the activation of hippocampal microglia and astrocytes, suggesting activation of an innate immune response in the brain. Interestingly, this activation persisted for two months post-exposure. Increased immunostaining for amyloid beta and a loss of synthapsin staining were also found. These data contribute to the growing evidence for an association between environmental exposures and neurodegenerative disease progression.

2482 Does Air Pollution Exposure Increase Risk for Autism?
I. Hertz-Picciotto1, H. E. Volk2 and R. S. McConnell3. "MIND Institute & Center for Children's Environmental Health, University of California Davis, Davis, CA; 2Center for Children’s Environmental Health, University of Southern California, Los Angeles, CA; Sponsor: D. Cory-Slechta.

Windham et al. (2006) reported that children with autism were more likely to live in census tracts with higher ambient levels of diesel, certain metals, and solvents. Traffic is a major source of air pollution, including particulates, polycyclic aromatic hydrocarbons, metals, sulfur dioxide and nitrogen oxides. Proximity to freeways and major roadways is highly correlated with ambient levels of such traffic-related air pollution. Using data from the CHARGE (CHildhood Autism Risk from Genes and Environment) Study, we examined the association between autism and prox- imity of residence to freeways and major roadways during pregnancy and near the time of delivery. (Volk et al 2011). Using data from 304 autism cases and 259 typi- cally developing controls, the mother’s address recorded on the birth certificate and trimester-specific addresses obtained by questionnaire were geocoded, and measures of distance to freeways and major roads calculated with ArcGIS software. After ad- justment for sociodemographic factors and maternal smoking, mothers of cases were more likely, at the time of delivery, to live within a quarter mile of a freeway as compared with mothers of controls: odds ratio (OR) = 1.86. At greater distances, risk was not elevated. Autism was also associated with residential proximity to a freeway during the third trimester (OR = 2.22). For each address, exposures to spe- cific air pollutants from traffic and stationary sources were estimated with U.S. EPA’s Air Quality System (Volk et al, in press). Exposures to nitrogen dioxide (NO2) and particulate matter less than 2.5 and 10 µm in diameter (PM2.5 and PM10) in the first year of life were associated with about a two-fold increased risk for autism (NO2 OR=2.06; PM2.5 OR=2.12; PM10 OR=2.14), with similar find- ings for gestational exposures. Given that many constituents in air pollution induce neurodevelopmental deficits in experimental rodent studies, a contribution from air pollution to autism susceptibility is plausible.

2483 Ozone Pollution, Oxidative Stress, and Dysregulation of Inflammatory Responses in Rat Hippocampus.

The oxidizing compounds contained in the air that we breathe in the large cities, is related to a large number of chronic degenerative diseases. To study the effects of ozone pollution on the brain, we developed a model consisting in a chronic expo- sure of rat to ozone doses, similar to total levels that occur in a day of high pollution in Mexico City. For this purpose, 84 male Wistar rats (250 to 300 g) were ran- domly divided into 7 groups, each group received one of the following treatments: 1) control, 2) exposed to ozone free air for 30 days, 3) exposed to ozone for 7 days, 4) exposed to ozone for 15 days, 5) exposed to ozone for 30 days, 6) exposed to ozone for 60 days, and 7) exposed to ozone for 90 days. (0.25 ppm /4 hours daily). Two hours after the last exposure to ozone, each group was randomly divided into
two subgroups, hippocampus were dissected and afterward processed for immunohistochemistry. The second subgroup was used for western blot and spectrophotometric techniques (to determine lipid peroxidation, oxidized proteins, superoxide dismutase and glutathione). The results indicate that ozone, causes an increase in lipids and proteins oxidation after 15 days of exposure. After 30 days we found in the hippocampus damage and decrease of the neuroblasts, and neurons number, glial activity, and increased of proinflammatory markers, together with changes in levels of antioxidant defenses and loss of its enzymatic activity. Results also showed accumulation of deposits of amyloid beta (Aβ) 1-42 at 90 days exposure. In conclusion: ozone lead to oxidative stress causing a neurodegenerative process in hippocampus, together with, dysregulation of immune responses, and loss of brain to capacity repair and Aβ 1-42 plaques formation at 90 day of ozone exposure. These experiments showed that chronic exposure to ozone at low doses, per se, is able to cause an oxidative stress state, which produces progressive cellular damage and cell death that resembling the damage described in the physiopathology of neurodegenerative diseases.

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2484 Translational Methods to Assess the Safety of Natural Health Products, Including Traditional Medicines and Dietary Supplements.

L. C. Griffiths and S. A. Jordan.

Globally, ~80% of the world’s population relies upon traditional medicines as part of standard healthcare; ~100 million Americans spend ~$28 billion annually to consume herbs, vitamins, minerals, amino acids, and other naturally occurring products in the form of dietary supplements, botanical drugs, and natural health products. The complexity of mixtures with variance in composition and quality presents a challenge for risk practitioners. To address these issues, new methodologies and predictive technologies are being developed, tested and validated to mitigate risk of human toxicity and effectively increase the quality of information useful for application in safety assessments. Many of these methods are already reflected in multiagency government initiatives. Such methods are referred to as “translational” for making use of and extrapolating data from scientifically defensible approaches towards establishing human safety and use of viable product. In this symposium, various in vitro, in vivo, state-of-the-art in silico (computational), and ‘omic methodologies will be presented. Discussion will cover case studies and regulatory science activities at US FDA/CBER and NCTR including conventional toxicological and computational assessments of individual chemical constituents and mixtures. Computational tools to deconvolute complexities and predict molecular targets for constituent phytochemicals will also be highlighted. Speakers will also address how the United States Pharmacopeia (USP) conducts evidence-based reviews on the safety of dietary supplements and how to couple data from in silico structure-based methods to compliment and strengthen evidence and support committee-based decision making regarding the use of dietary supplements for monograph development on product quality. Collectively, these presentations will provide a global picture of the state and utility of modern methods for assessing toxicity and safety of these products.

2485 Session Overview: The Promise of Translational and Integrative Safety Assessments of Dietary Supplements, Traditional Medicines, and Herbal Drugs.

S. A. Jordan.

For thousands of years, herbs and other natural substances have been used in health care. Some traditional healing paradigms include knowledge of potential toxicity, and ways of preventing adverse effects. However, with modern use, a much larger and varied population now uses these products. In addition, commercial products may not be true to traditional use, indication, or form. A rise in use of these products with pharmaceutical drugs, resulting in possible herb-drug interactions, also has become an issue of concern. Potential safety concerns may also arise from quality issues. Regulators are often faced with a lack of quality information when making risk-based decisions on the safety of dietary supplements or traditional medicines, and an increase in these data allows for improved decision making. The development of translational methods, including omics, and computational models, has presented an opportunity to increase the amount of information, and to integrate novel data with traditional methods of assessing toxicity. The integration of all available data, including those derived from translational studies, can reduce uncertainty in decision making related to the potential hazard and risk associated with the use of these product types. Overall, this will benefit the public, the industry, and the regulatory community which is charged with licensing and monitoring the safety of dietary supplements and traditional medicines.

2486 In Silico Methods As Translational Tools for Supporting the Safety Assessment of Natural Health Products.

L. G. Valerio.

In silico methods including computational toxicology can serve to help address data gaps during safety evaluations for regulatory and industrial product safety. Natural health products including dietary supplements, botanicals, herbs, and related substances can benefit from the wide spectrum of scientific evidence produced by in silico toxicology analyses. Discussion will cover case studies and regulatory science activities in the application of structure-based computational assessments of individual chemical constituents derived from natural products and mixtures modeling. Translation of these data to support human safety and risk assessment processes will be presented. The in silico approach as a translational tool with emerging evidence-based predictive technologies and the use of these methods in applied safety science is of extraordinary heightened interest with the goal of meeting today's needs for protecting public health.

2487 In Vitro and In Vivo Approaches for Assessing the Safety of Natural Health Products.

W. E. Salminen.

Dietary supplement use continues to increase as many people see these “natural” products as inherently safe alternatives to drugs. However, in the US and many other parts of the world, dietary supplements require none to minimal safety data before they are marketed. Many dietary supplements are complex mixtures of a wide variety of chemical compounds and the composition can vary greatly from manufacturer to manufacturer and even batch to batch making safety assessments very difficult. Various in vitro and in vivo approaches for assessing the safety of dietary supplements from classical toxicology studies to state-of-the-art toxicogenomics assessments will be reviewed. An overview of the US National Toxicology Program’s (NTP) testing of dietary supplements will be presented and examples of potential drug-dietary supplement interactions will be discussed since these are likely to increase as people use dietary supplements in combination with approved drugs.

2488 Computational Methods Linking Traditional Chinese Medicine (TCM) and Western Therapeutics.

D. E. Johnson.

Herbal remedies are widely used throughout the world with approximately 80% of the global population relying on traditional medicines as part of standard healthcare. In the US, an estimated 1 in 5 adults regularly consume herbal products and most of these are not included in patient records as “other medications”, thereby limiting the understanding of potential herb-drug interactions to both the patient and health care professional. We have been using computational tools to deconvolute complex recipes and predict molecular targets for constituent phytochemicals with the goal of forming a comprehensive tool to help predict herb-drug interactions. This work will be illustrated using both open-source tools and commercial biological pathway mapping and predictive algorithms. Translational methodology will be introduced that proposes a chemical-disease category linkage between TCM and Western medicine for a rational integrative medicine approach.


Y. Kang.

Natural health products, including traditional Chinese medicines (TCMs) are often complex mixtures. Assessing the safety of individual components of such products is problematic. The use of TCM has raised concerns of heavy metal contamination and toxicity. However, it has been known that metals and metalloids are essential components in some TCM preparations. For instance, mercury in cinnabar is irreplaceable for its therapeutic effect. Another scenario is that the TCM formulation is toxic to healthy population, but becomes remedy to seriously sick people. In this context, the dose regimen of arsenic trioxide used to treat acute promyelocytic leukemia (APL) is extremely toxic to general population, but is of high therapeutic efficacy for APL patients. It is difficult, if not impossible, to distinguish metal composition from metal contamination in TCM and their associated therapeutic or toxic effects. A novel approach of herbogenomics can provide
alternate perspectives on the active ingredients and toxic transformations of TCM. The overall effects of TCM on target organs can be elucidated by functional genomics and proteomics, thus the addition or subtraction of heavy metals from TCM preparations would affect the outcome, which can be reflected by the genomic profile alterations. The information about metal composition and contamination is critical for general population considering the use of TCM as an alternative remedy.

2490 Evidence-Based Reviews As a Method for Assessing the Safety of Dietary Supplements.

M. L. Hardy. Center for Integrative Medicine, University of California Los Angeles, Los Angeles, CA. Sponsor: J. Griffiths.

Dietary supplement safety is one of the main regulatory concerns in the United States. Adverse events occur for a variety of reasons including contamination, misidentification or adulteration of plant based dietary supplements. However, even when a product is correctly made, adverse events in human subjects have been reported. Systematic review and meta-analysis of published clinical trials and adverse event case reports allow aggregation of disparate reports in order to improve the analytic strength of evaluations of potential harms. When adverse events are reported in properly randomized clinical trials, causality of the reported adverse event can be assumed. Clinical trials are almost always of insufficient power to detect any but the most common adverse events. Rare or serious adverse events are usually reported in as case reports or small case series. Attribution of risk in this case is limited by a number of factors including timing and duration of exposure to suspected offending substance as well as the clinical circumstances of the user. Although causality can rarely be attributed based on case reports, systematic review of aggregated cases can identify a suspicious signal. Additional limitations of all sources of adverse event reports are the result of incomplete information including very often poor characterization of the potential offending material. The application of these principles will be illustrated using an analysis of adverse events related to ephedra.

2491 Are We Like Rodents, Rabbits, or Something Else? Mechanisms of Developmental and Reproductive Toxicity across Species.

R. J. Rasoulipour1, K. Johnson1, S. N. Campion2, C. Timchalk1 and J. E. Goodman1. 1The Dow Chemical Company, Midland, MI; 2Pfizer Global, Groton, CT.

Given the inherent complexity of embryo/fetal development and reproductive biology, developmental and reproductive toxicity (DART) hazard identification and research still heavily relies on animal models. Typically, this type of research is conducted in rodents (e.g., mouse and rat) as well as nonrodent species (e.g., rabbit and nonhuman primate), which, in the face of toxicity findings, raises the question of relevance to humans. Are we more like mice, rats, or other model organisms? Not surprisingly, answering this question is a challenge, and the scientific approach may be quite different depending upon the biological system and the level of mechanistic information available. However, within this challenge is an opportunity to understand the toxicokinetic and toxicodynamic differences between the species and provide the appropriate context for developmental and reproductive findings. Providing this context can directly impact the risk assessment and regulatory decision-making process. This workshop will highlight several different approaches to relate animal reproductive and developmental toxicity findings to human health. Speakers in fields of basic research, product safety testing, epidemiology, and physiologically-based pharmacokinetic modeling will address the central theme of the workshop, which is applying different experimental strategies to analyze cross-species toxicity. Within each presentation, potential regulatory implications and feedback from agencies when available, will be addressed. The ultimate goal of the workshop are an increased understanding of the different approaches to dissect complex toxicity mechanisms and to take the next step towards cross-species comparisons that impact human relevance and regulatory decisions.


R. J. Rasoulipour. Developmental and Reproductive Toxicology, The Dow Chemical Company, Midland, MI.

This presentation will provide a case study on unique challenges often posed by elucidating developmental toxicity mechanisms identified within guideline toxicity studies. The case study will focus on a recently identified novel mechanism of rat-specific developmental toxicity induced by a developmental molecule, sulfoxafolt. At high doses, rats exhibited ear and limb flexure effects occurred in rats, but not rabbits. The proposed mode-of-action was that these effects had a single mechanism mediated via the rat fetal isoform of the muscle nicotinic acetylcholine receptor. The studies included a combination of novel in vivo and in vitro mechanistic studies, which were integrated to identify agonism on this receptor as a critical event. Moreover, species comparison studies revealed that this initial event was isolated to sustained agonism on the rat fetal isoform of the nicotinic acetylcholine muscle receptor and did not occur in humans. Feedback on this project from global regulatory agencies will be presented and discussed. This stage-setting talk provides a clear example of how human relevance can be understood if the toxicity is mediated by a relatively simple mechanism (e.g., agonism at a single receptor).
The developing brain is vulnerable to insults and evidence implicates low-level chemical exposures as potential developmental neurotoxins. Organophosphorus (OP) insecticides, like chlorpyrifos (CPF), inhibit cholinesterase (ChE) and are neurotoxic. PBPK/PD models have been exploited to enable cross-species extrapolation of CPF brain dosimetry and ChE inhibition and when linked with a dietary exposure model can predict response across populations. However, recent epidemiology studies suggest that OP neurotoxicity occurs at low-doses, even in the absence of significant brain ChE inhibition. The lack of quantitative cross-species brain dosimetry data associated with epidemiology results hampers any mechanistic based risk assessments. The implication of localized heterogeneous CYP450 brain metabolism has historically not been extensively investigated, but recent research suggests it is of key importance. To address these limitations, we are testing the hypothesis that low-dose exposures of preweanling rats to OP insecticides will result in differential brain region dosimetry, enhanced by localized bioactivation, potentially resulting in subtle changes in brain chemistry. Comparative in vitro metabolism studies in rats indicate that the overall brain microsomal metabolism was a fraction (~3%) of the liver. Following in vivo administration (1 and 5 mg/kg/day) of CPF to post-natal day-10 pups, CPF and its major metabolite were quantified in the brain with evidence of regional deposition and localized metabolism. The importance of localized brain metabolism is highly relevant for lipophilic pesticides that sequester in the lipid rich regions of the brain and can undergo local metabolic activation to produce neurotoxic effects. This is particularly important in juvenile animals, and children, where there may be a disproportionate deposition of the parent pesticide in the brain. In this regard, these PBPK/PD modeling strategy have significant regulatory implications for assessing developmental neurotoxicity.

Using Epidemiology to Analyze Neurodevelopmental Toxicity across Species.

J. E. Goodman, Gradient, Cambridge, MA.

The final presentation of the session will tackle cross-species analysis from a different angle by utilizing epidemiology data. This presentation will explore how epidemiology studies can address toxicity across species by estimating human exposures and/or outcomes with biomarkers and putting these into context with the animal model results. It will focus on how the effect of timing, selection choice, and measurement of biomarkers, as well as the results of toxicity studies, can influence the interpretation of results. Chlorpyrifos and neurodevelopmental effects were used as a case study. US EPA assessed whether epidemiology data suggest that fetal or early-life chlorpyrifos exposure causes neurodevelopmental effects and, if so, whether they occur at exposures below those causing 10% inhibition of blood acetylcholinesterase (AChE), which is currently considered the most sensitive endpoint. We conducted a hypothesis-based weight-of-evidence analysis and found that a proposed causal association between chlorpyrifos exposure and neurodevelopmental effects in the absence of AChE inhibition does not have a substantial basis in existing animal or in vitro studies, and there is no plausible basis for invoking such effects in humans at their far lower exposure levels. The epidemiology studies fail to show consistent patterns; the few associations are likely attributable to alternative explanations. The human data are inappropriate for a dose-response assessment because biomarkers were only measured at one time point, may reflect exposure to other pesticides, and many values are at or below limits of quantification. When considered with pharmacokinetic data, however, these biomarkers provide information on exposure levels relative to those in experimental studies and indicate a margin of exposure of at least 1,000. Because animal data take into account the most sensitive lifetimes, the use of AChE inhibition as a regulatory endpoint is protective of adverse effects in sensitive populations.

Cumulative Risk: Toxicity and Interactions of Physical and Chemical Stressors.


Recent efforts to update cumulative risk assessment procedures and develop community-based risk assessment methods reflect increased interest in incorporating the totality of variables affecting human health into the risk assessment process. One key roadblock in advancement is uncertainty as to how nonchemical stressors behave in relationship to chemical stressors. An assumption that simplifies incorporation of nonchemical stressors into current risk assessment paradigms is that nonchemical stressors act in the same manner as chemicals. However, evidence is required to support this assumption. The term nonchemical stressors encompasses a diverse set of variables including physical stressors, such as noise, temperature, disease, and pollution, as well as psychosocial stressors, which involve perception of circumstances. Physical stressors offer a reasonable starting point for measuring the effects of nonchemical stressors and their modulation of chemical effects (and vice versa), as they clearly differ from chemical stressors, present many diverse and highly-relevant stressors, and “doses” of many physical stressors are easily quantifiable. The relative importance of nonchemical stressors and their modulation of the impact of nonchemical stressors on chemical-mediated toxicity or the joint impact of coexposure to chemical and nonchemical stressors. While generally true, there are several instances where a substantial body of evidence exists. The objective is to provide expert overview, for those chemical and physical stressors that have been sufficiently studied to get at least a limited understanding of their joint impact. In addition to providing the current state of knowledge, data gaps will be identified that should be addressed to facilitate inclusion of nonchemical stressors in risk assessment. (This abstract does not reflect US EPA or NIEHS policy.)

Cumulative Risk: Chemicals and Infectious Disease.

M. Selgrade, ICF International, Durham, NC.

At least 4 types of mechanisms underlie potential interactions between toxic chemicals and infectious disease. 1) The best understood is suppression of immune responses, resulting in increased incidence/severity of infectious disease. For example, decreased alveolar macrophage function following exposure to several air pollutants enhances the risk of certain bacterial infections. Research on this model provides both qualitative and quantitative approaches to describe this risk. 2) Certain immunomodulatory mediators that are activated during chemical exposure and include cytokines and immunometabolic enzymes and transporters and have the potential to alter chemical toxicity as illustrated by the effects of murine cytomegalovirus on parathion poisoning, sodium pentobarbital induced sleeping time, and cyt P450. Infection and inflammation have also been shown clinically to affect the metabolism, distribution, and elimination of certain drugs. 3) Chemical exposure may enhance inflammation and immune pathology associated with an infection. This is best illustrated by effects of ozone, ultraviolet radiation, and TCDD on influenza infection. In all cases mortality is enhanced in the absence of increased virus titers in the lung or viral dissemination. Deaths appear to be due to increased inflammatory responses. Similarities exist between receptors and subsequently triggered signaling pathways for pathogen associated molecular patterns (PAMPs) and damage associated molecular patterns (DAMPs), which trigger inflammatory responses. A systems approach that examines the integration of these pathways is needed to better describe this phenomenon. 4) Infection enhances chemical induced lesions, e.g., p53 mutations, inflammation, cell proliferation. Such mechanisms might explain the interaction between hepatitis B virus infection and aflatoxin in the induction of liver cancer. These mechanisms are not necessarily comprehensive, distinct, or mutually exclusive and are imperfectly understood. However, they provide a useful framework to further explore the interactions between infectious disease and exposure to toxic chemicals with the ultimate goal of improving our understanding of cumulative risk.

Enhancement of Noise-Induced Hearing Loss by Chemicals.

Y. C. Mozata, Division of Applied Research & Technology, NIOSH, Cincinnati, OH. Sponsor: J. Simmons.

The auditory effects of chemical toxicants have been investigated in the past two decades, in animal and human field and clinical studies. A number of studies demonstrated that some solvents, metals, asphyxiants, pesticides not only affect the sensory organ of the auditory system, as noise does, but also affect central auditory structures. Ototoxicity induces outer hair cell dysfunction in the cochlea (similar to the effects of noise), whereas neurotoxicity induces central auditory dysfunction. Audiological signs of neurotoxicity may or may not include poorer hearing thresholds, in addition to difficulties discriminating sounds such as speech, particularly in adverse listening conditions. The existing evidence prompted the proposal of new guidelines and standards on hearing loss prevention. In the U.S., the National Institute for Occupational Safety and Health has discussed specific research needs regarding the ototoxicity of chemicals used at work. The American Conference of Governmental Industrial Hygienists and the U.S. Army have proposed preliminary practical steps that employers and occupational health professionals can take to improve hearing loss prevention. Australia and New Zealand have developed standards recommending hearing tests for workers exposed to ototoxic agents. In the legislative arena, the European Parliament published a new noise directive (2003/10/EC), that requires employers to give attention to any effects on workers’ health and safety resulting from interactions between noise and work-related ototoxic substances, when performing risk assessment of workplaces. Legislation regarding compensation has also changed in Australia and Brazil. In this presentation
the auditory effects of chemical alone or in combination with noise will be reviewed, and the recent public debates, legislative developments and alternative strategies for the prevention of auditory effects resulting from exposure to chemicals in the workplace will be presented.

2500 Exacerbation of Toxicity of Air Pollutants and Pesticides by Thermal Stress.

Considering the likelihood of global warming in the near future, it is important to understand how heat stress will alter the health effects of toxicants. The toxicity of pesticides and airborne toxicants is generally exacerbated in a warm environment. As air temperature increases, the pulmonary intake of air pollutants and absorption of pesticides applied to the skin is generally accelerated. Cellular toxicity is typically exacerbated when body temperature is elevated. This is primarily a result of a Q10 effect, meaning that the rate of a biochemical reaction doubles with a 10 °C increase in tissue temperature. The generation of reactive oxygen species is also exacerbated with warmer temperatures.

Since warmer temperatures worsen chemical toxicity, a thermoregulatory response to lower body temperature can be protective. Hyperthermia is most likely going to be detrimental in the recovery from toxicant exposure. Rodents and other small mammals have relatively large surface area:volume ratio. Following exposure to pesticides and air pollutants, metabolism is reduced and a rapid reduction in body temperature ensues. If given the opportunity to thermoregulate behaviorally, rodents seek colder temperatures, allowing body temperature to decrease quickly. Thus, the hyperthermic response is a regulated response. This is thought to be an adaptive response because the hyperthermic response is protective. Large mammals, including humans, have a greater thermal inertia and are unable to mount a hyperthermic response as is seen in rodents. A warmer environmental temperature will thus impede the hyperthermic response to toxicant exposure in rodents and will also be stressful to large mammals that are unable to undergo a significant cooling response. With the potential impact of global climate change on increased incident of heat stress in urban areas that are also rife with pollution, the topic of the thermoregulatory responses to environmental toxicants is timely. This is an abstract of a proposed presentation and does not reflect US EPA policy.

2501 Modulation of X-Ray Mediated Testicular Toxicity by Chemical Exposure.
K. Boekelheide, Brown University, Providence, RI.

High density microarrays and a detailed bioinformatics analytical approach were used to demonstrate that an initial chemical exposure to 2,5-hexanedione (HD) altered the rat testis to ameliorate the response to a subsequent exposure to x-irradiation. Adult male rats were exposed to HD (0.33% or 1%) in the drinking water for 18 days followed by x-ray (2Gy or 5Gy), resulting in a total of 9 treatment groups. Testis samples were collected after 3 hr and gene array analysis was performed. Using a novel bioinformatic approach to summarize the effect of HD across all treatment groups, we focused on the modulation of x-ray-induced gene alterations by HD co-exposure. Enrichment analysis was used to identify biological pathways where HD modification of gene expression was the greatest. HD exerted a significant influence on genes involved in Cell Cycle and DNA Replication, Recombination, and Repair. HD also had an antagonistic effect on x-ray-induced alterations of several apoptotic genes (Fas, BBC3, AEN). To further investigate the specific cell populations and stages in which these critical gene alterations occur, laser capture microdissection (LCM) samples were collected from the basal compartment of the seminiferous epithelium, enriching for those germ cells most susceptible to x-ray-induced apoptosis. Quantitative RT-PCR of the LCM samples confirmed the suppression of apoptotic genes by HD co-exposure. The co-exposure attenuation of germ cell apoptosis is the result of an adaptive response to the chemical exposure, causing altered paracrine signalling of the supportive cells in the seminiferous epithelium. These results suggest that toxicity pathway responses determine the outcome of co-exposures, whether chemical or physical in nature, and that complex paracrine interactions between cells modulate the extent of injury.

2502 Sunlight Enhancement of the Toxicity of Air Pollutant Mixtures.
K. G. Sexton1, J. Zavala1, B. O’Brien1, W. Vizuete1, R. C. Fry1, J. Laspesa1,2 and J. Rusyn1. 1Environmental Sciences & Engineering, University of North Carolina at Chapel Hill, Chapel Hill, NC; 2Center for Environmental Medicine and Lung Biology, University of North Carolina at Chapel Hill, Chapel Hill, NC.

Sunlight can significantly drive photochemical reactions of mixtures of air pollutants commonly observed in the atmosphere, producing many well known toxic compounds such as formaldehyde and other carbonyl containing products. These reactions also contribute to the formation of secondary organic aerosols as well as modifying the composition of existing aerosols or particulate matter (PM). Smog chambers can be used to prepare repeatable, controlled mixtures of simple to increasing complexity and be used to study photochemical atmospheric transformation with natural sunlight or simulated sunlight. Smog chambers can be interfaced with direct exposure to in vitro or in vivo models for toxicity studies including direct air-liquid-interface in vitro or in vivo inhalation exposures. Photochemical experiments have been conducted in smog chambers with industrial mixtures and complex mixtures of motor vehicle exhaust in urban atmospheres, often demonstrating enhanced toxicity as measured by markers of inflammation and other biological endpoints such as cytotoxicity. Modifications of experiments, exposure conditions and additional toxicological analyses can provide mechanistic and mode of action understanding. Novel genomic analyses of cells exposed to an urban-like mixtures showed transcriptional changes on a subset of genes, increasing the number of genes with altered expressions from 19 for the un-irradiated mixture, to 709 genes after a one-day sunlight irradiation. The implication is that toxicologists and those dependent on their findings, should consider studies that include mixtures resulting from natural atmospheric photochemical reactivity, transformation of air components, and the resulting enhancement of toxic effects of air pollutants and their products. Not considering such effects, could result in misinterpreting the mode of action and underestimating the potential risk of exposure to air pollution mixtures or its sources.

2503 Mechanistic, Occupational, and Clinical Aspects of Lead Exposure.
A. Vale. School of Biosciences, University of Birmingham, Birmingham, United Kingdom.

The mechanisms of lead toxicity are increasingly being explained by the ubiquitous reactivity of the bivalent lead cation and its ability to substitute for essential cations, notably calcium and zinc. By these means, lead complexes with functional groups including thiol and carbonyl groups, and damages many fundamental cell processes and structures including enzyme pathways, phospholipid integrity, ion channel specificity and control, and intrinsic protective systems including free radical scavengers and cellular repair mechanisms. Owing to the large sample sizes involved and its nationally-representative nature, NHANES has been the subject of a number of epidemiological analyses relating blood lead concentrations to a range of adverse outcomes such as blood pressure, renal function, auditory thresholds, and a host of other cardiovascular, neurobehavioral, and other developmental or adult outcomes. Some practitioners are now proposing that, as the NHANES data suggest that lead concentrations even less than 5 μg/dL (0.24 μmol/L) can have some health consequences, chelation should be performed at even very low lead concentrations. Is this an appropriate interpretation of these data? There is concern that the occupational intervention concentrations worldwide are not only unsupported scientifically and clinically but also have been set at concentrations that permit unsafe practices to continue. A group of experts has proposed that workers should be removed from occupational exposure if a single blood lead concentration exceeds 30 μg/dL (1.45 μmol/L), or if two successive blood lead concentrations measured over a four-week interval equal or exceed 20 μg/dL (0.97 μmol/L). Will these recommendations prevent clinically significant occupational lead exposure? Due to the paucity of clinical data, there is controversy about the lead concentration at which chelation therapy should be instituted in adults when exposure prevention has failed, the antidote to be used, and the most effective regimen to be employed.

2504 Novel Mechanisms of Toxicity.
B. Lantz. Cellular and Molecular Medicine, University of Arizona, Tucson, AZ.

The broad spectrum of lead toxicity with adverse manifestations in developmental and functional aspects of many if not all organ systems are increasingly being explained by the ubiquitous reactivity of the bivalent lead cation and its ability to substitute for essential cations, notably calcium and zinc. By these means lead com-
plexes with important functional groups including thiol and carboxyl groups and damages many fundamental cell processes and structures including enzyme pathways, phospholipid integrity, ion channel specificity and control and intrinsic protective systems including free radical scavengers and cellular repair mechanisms. The ability for lead to substitute for calcium causes erroneous activation of calcium-dependent proteins and modulation of calcium-sensitive receptors. At its extreme, impaired regulation of calcium transport results in intracellular calcium accumulation which triggers apoptosis. Lead can activate protein kinase C to disrupt intracellular regulatory processes causing incorrect gene expression resulting in disordered cell proliferation and differentiation. Lead-mediated malfunction of the calcium sensitive N-methyl-D-aspartate (NMDA) receptor is recognized as one of, if not the most, important mechanism of lead-induced damage to neuronal development, learning and memory. As the details of the molecular mechanisms of lead toxicity are unravelled, it is becoming clear that their complexity is increased by the fact that lead, like many toxins, acts not in isolation but as part of a multifactorial armoury of adverse influences including environmental factors, which together dictate the development, manifestations and progress of disease.

The National Health and Nutrition Examination Survey (NHANES) are cross-sectional surveys of the civilian noninstitutionalized population of the United States that have been administered by the National Center for Health Statistics and ongoing for decades. Subjects are selected based on a stratified multistage probability sampling of counties, blocks, households, and persons within households, with oversampling of some population subgroups (such as Mexican Americans, non-Hispanic blacks, and adults 60 years or older). Evaluations included the administration of extensive questionnaires, physical examination, and collection of urine and venous blood for a wide range of laboratory analyses. Lead levels have been measured in samples of venous blood since the 1970s. Owing to the large sample sizes involved (typically >10,000 individuals) and its nationally-representative nature, NHANES has been the subject of a number of epidemiologic analyses relating blood lead levels to a range of outcomes such as blood pressure, renal function, auditory thresholds, and a host of other cardiovascular, neurobehavioral and other developmental or adult outcomes. This presentation will provide an overview of these studies from the perspective of understanding dose-response relationships; it will also compare them with the parallel body of recent epidemiologic studies using other biomarkers of lead exposure (e.g., XRF-measured bone lead levels as a marker of cumulative exposure) and/or other study designs (e.g., prospective or case-control studies).

Research findings have heightened public health concerns regarding the hazards of low dose lead exposure to adults and children. In adults, studies have established the potential for hypertension, effects on renal function, cognitive dysfunction and adverse female reproductive outcome in adults with whole blood lead concentrations less than 40 μg/dL (1.93 μmol/L). However, in most nations worldwide, regulatory occupational exposure limits permit workers to maintain blood lead concentrations in excess of 40 μg/dL for a working lifetime. A group of experts has recently recommended that workers undergo removal from occupational lead exposure if a single blood lead concentration exceeds 30 μg/dL (1.45 μmol/L), or if two successive blood lead concentrations measured over a four week interval equal or exceed 20 μg/dL (0.97 μmol/L). Removal from lead exposure should be considered to avoid long-term risk to health if exposure control measures over an extended period do not decrease blood lead concentrations below 10 μg/dL (0.48 μmol/L), or if selected medical conditions exist that would increase the risk of continued exposure. In order to assure reductions in permissible blood lead concentrations, medical surveillance for lead exposed workers is recommended to include quarterly blood lead measurements on individuals with blood lead concentrations between 10 to 19 μg/dL (0.48 – 0.92 μmol/L), and semi-annual blood lead measurements when sustained blood lead concentrations are less than 10 μg/dL (0.48 μmol/L).

Intravenous edetate calcium disodium and oral succimer (dimercaptosuccinic acid; DMSA) are potent chelators of lead approved in many countries for the clinical management of lead poisoning. Both agents reduce the body burden of lead by the renal elimination of a lead-chelating agent complex but there remain unanswered questions regarding their precise pharmacokinetic and pharmacodynamic properties. These include the specific effects of each agent on tissue lead distribution and mobilization and how these parameters are affected by dose and duration of both lead poisoning and chelation. Edetate calcium disodium chelates by exchanging its central calcium ion for lead but the exact chemical nature of the succimer-lead chelate in man remains ill-defined. While assessment of efficacy of both agents is hampered by limited data, tissue lead concentration, bioavailability of lead toxicity, variations in study design, species differences and, for clinical studies, the subjective nature of reporting changes in symptoms, available data overall suggest both drugs offer similar efficacy with regard to enhancing lead elimination, though this remains controversial. In addition, the indications for chelation in man remain ill-defined. The adequacy of treatment remains contentious both because of the paucity of efficacy data and because of increasing evidence that toxic effects of lead sustained during early development cannot be reversed by chelation. Moreover, even though most clinical toxicologists agree that where exposure prevention has failed, chelation should be considered in adults whose blood lead concentrations ≥ 50 μg/dL (2.4 μmol/L), the most effective regimen to employ in these circumstances is still controversial.

The eye is a unique organ composed of many different structures working together to facilitate vision. The maintenance of clear vision during aging is threatened by various physiological changes (e.g., presbyopia) or diseases (e.g., age-related macular degeneration). The need for treatments is of increasing importance as the size of the aging population grows. The National Eye Institute predicts that by 2020, more than 50 million Americans will be impacted by age-related eye disease. Some of these conditions have been overcome through the use of medical devices. Ocular medical devices consist of instruments, apparatuses, appliances, and materials. Some devices are purely structural; whereas, other medical devices are a part of a delivery system that releases a drug. The five presentations in this program are designed to educate the audience on the therapeutic, safety, and regulatory challenges of developing ocular medical devices and drug delivery systems. The first presentation will cover the special requirements for developing contact lenses and contact lens solutions. The second presentation will describe the challenges associated with accommodating intraocular lenses (IOLs). The third presentation will discuss the ocular barriers (e.g., blood-eye-barrier) and how ocular medical devices and sustained release drug formulations have been designed to address these barriers with respect to developing protein therapeutics. Development of and regulatory challenges associated with a unique biodegradable small molecule ocular drug/injection applicator delivery system will be discussed in the fourth presentation. The symposium will conclude with a presentation that discusses the safety and regulatory requirements and complexities pertaining to ocular medical devices and ocular drug delivery systems.
biocompatibility testing strategies for contact lenses and solutions. For example, depending on the type of assay, the tested material may need to be the formulation itself, a lens extract or a lens/formulation combination. Further, it has been demonstrated that the selection of assays and individual study designs can influence the outcome of the study, which could potentially impact product registration. Testing protocols therefore need to be designed to account for the unique chemical and physical properties of the ocular medical device being tested in order to avoid potentially false positive outcomes. Consequently, a rational science-based approach should be used to develop and justify the biocompatibility testing strategy and study designs to ensure that the biological evaluation conducted is appropriate, robust and Regulatory-acceptable. This presentation will highlight some of the key challenges and case studies when conducting biocompatibility evaluations on contact lenses and solutions.

2510 Accommodative Intraocular Lenses: A New Class of IOLs with a New Class of Challenges.

A. Glasser, College of Optometry, University of Houston, Houston, TX. Sponsor: L. Render.

A new class of intraocular lenses (IOLs) is being developed with the goal of restoring the focusing ability of the eye (accommodation) for treating presbyopia (the age-related loss of near focusing ability). Although cataract surgery may be among the safest and most common of surgical procedures, these so called accommodative IOLs (A-IOLs) present new challenges as well as offer new opportunities. A-IOLs differ considerably from standard IOLs in that they are biomechanical devices designed to move or change shape in response to ciliary muscle contraction. A-IOLs are bulkier, made from different materials, may be implanted in different locations in the eye and have mechanisms of action that differ fundamentally from standard IOLs. The surgical procedures required to implant these devices are also more challenging, as they require new surgical devices. In addition, the safety considerations for A-IOLs are different. Pre-clinical animal testing that can be done is limited, complications that can arise are unique, and regulatory hurdles for demonstrating safety and effectiveness are higher. The ultimate success of A-IOLs will rely on solving significant biological challenges that still remain, such as resolving the post-operative healing response of the eye including prevention of post-operative lens epithelial cell proliferation and fibrosis of the lens capsule. This presentation will describe the challenges associated with the development A-IOLs. It will also introduce some future potential applications such as, the possibility of delivering drugs from A-IOLs to solve these biological challenges, as well as other unique surgical and pharmacological interventions aimed at resolving the problem of presbyopia.

2511 Safety Assessment Strategies and Challenges for Developing Intravitreally Administered Biologics.

E. A. Thackaberry, Safety Assessment, Genentech, Inc, San Francisco, CA.

The development of intraocular drugs to treat posterior segment disease, such as age-related macular degeneration, presents both advantages and challenges due to the unique aspects of ocular anatomy and physiology. While drug administration within the eye affords the promise of direct and local delivery, the reduction of systemic exposure and toxicity, and the availability of non-invasive tools allowing for real-time monitoring of the eye, there is an unmet need to reduce intravitreal injection frequency and treatment burden through sustained delivery formulations and devices. Adding to these complexities, developing antibody-based therapeutics presents additional challenges due to their inherent specificity (often limited to pri-mates), which may limit the animal models available for assessing safety. Case studies on the safety assessment strategies and challenges encountered in developing protein therapeutics to the posterior segment of eye will be presented.

2512 Development and Regulatory Considerations for an Ocular Drug Release System for Ocular Disease.

A. Wiese, Nonclinical Safety, Allergan, Irvine, CA.

Delivering drugs to various compartments of the eye presents unique scientific and regulatory challenges. This presentation describes these challenges in the context of the development of Ozurdex®, a bioerodable drug delivery system containing dexamethasone delivered to the posterior segment of the eye. Key to the development of this drug was understanding the drug distribution and its effects in the eye, drug release characteristics and erosion profile of the implant, and evaluation of the toxicological profile of both drug and implant. Additionally, and in parallel, development and testing of the applicator and consideration of drug interactions was ongoing. Regional differences in regulatory agency expectations will be presented. Finally, general learnings and applicability of those learnings to the development of drug delivery systems to various compartments of the eye will be discussed.

2513 Regulatory Considerations in Ocular Medical Device and Drug Delivery Systems Development.

C. Ghosh, Center for Drug Evaluation Research, US FDA, Silver Springs, MD.

Ocular medical devices and drug delivery systems encompass a wide variety of products including solid devices (such as intraocular lenses, surgical instruments and contact lenses) and devices that are liquid-based (such as viscoelastics and contact lens solutions). This presentation will present the classification and regulatory requirements for major categories of ocular medical devices and drug delivery systems and discuss the regulatory decision making and implications pertaining to preclinical and clinical evaluations.
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