Bioinformatics, 39(1), 2023, btac791 https://doi.org/10.1093/bioinformatics/btac791 Advance Access Publication Date: 8 December 2022 Applications Note



Data and text mining

SysBioIPGWAS: simplifying post-GWAS analysis through the use of computational technologies and integration of diverse omics datasets

Oluwadamilare Falola (1) 1^{1†}, Yagoub Adam (1) 1[†], Olabode Ajayi³, Judit Kumuthini³, Suraju Adewale¹, Abayomi Mosaku¹, Chaimae Samtal⁴, Glory Adebayo^{1,5}, Jerry Emmanuel^{1,2}, Milaine S. S. Tchamga⁶, Udochukwu Erondu⁷, Adebayo Nehemiah (1) 7, Suraj Rasaq⁷, Mary Ajayi⁷, Bola Akanle^{1,8,9}, Olaleye Oladipo^{1,8,9}, Itunuoluwa Isewon^{1,2,9}, Marion Adebiyi^{1,7,9}, Jelili Oyelade (1) 1,2,9 and Ezekiel Adebiyi (1) 1,2,9,10 **

¹Covenant University Bioinformatics Research (CUBRe), Covenant University, Ota, Ogun State 112104, Nigeria, ²Department of Computer & Information Sciences, Covenant University, Ota, Ogun State 112104, Nigeria, ³South African National Bioinformatics Institute, Life Sciences Building, University of Western Cape, Cape Town 7535, Republic of South Africa, ⁴Laboratory of Biotechnology, Environment, Agri-food and Health, Faculty of Sciences Dhar El Mahraz, Sidi Mohammed Ben Abdellah University, Fez 30000, Morocco, ⁵Department of Biological Sciences, Covenant University, Ota, Ogun State 112104, Nigeria, ⁶African Institute for Mathematical Sciences (AIMS), Muizenberg, Cape Town 7945, South Africa, ⁷Department of Computer Science, Landmark University, Omu-Aran, Kwara State 251103, Nigeria, ⁸Center for System and Information Services, Covenant University, Ota, Ogun State 112104, Nigeria, ⁹Covenant Applied Informatics and Communication Africa Center of Excellence (CApIC-ACE), Covenant University, Ota, Ogun State 112104, Nigeria and ¹⁰Applied Bioinformatics Division, German Cancer Research Center (DKFZ), Heidelberg 69120, Germany.

Received on June 23, 2022; revised on October 28, 2022; editorial decision on December 3, 2022; accepted on December 7, 2022

Abstract

Motivation: Post-genome-wide association studies (pGWAS) analysis is designed to decipher the functional consequences of significant single-nucleotide polymorphisms (SNPs) in the era of GWAS. This can be translated into research insights and clinical benefits such as the effectiveness of strategies for disease screening, treatment and prevention. However, the setup of pGWAS (pGWAS) tools can be quite complicated, and it mostly requires big data. The challenge however is, scientists are required to have sufficient experience with several of these technically complex and complicated tools in order to complete the pGWAS analysis.

Results: We present SysBiolPGWAS, a pGWAS web application that provides a comprehensive functionality for biologists and non-bioinformaticians to conduct several pGWAS analyses to overcome the above challenges. It provides unique functionalities for analysis involving multi-omics datasets and visualization using various bioinformatics tools. SysBiolPGWAS provides access to individual pGWAS tools and a novel custom pGWAS pipeline that integrates several individual pGWAS tools and data. The SysBiolPGWAS app was developed to be a one-stop shop for pGWAS analysis. It targets researchers in the area of the human genome and performs its analysis mainly in the autosomal chromosomes.

Availability and implementation: SysBiolPGWAS web app was developed using JavaScript/TypeScript web frameworks and is available at: https://spgwas.waslitbre.org/. All codes are available in this GitHub repository https://github.com/covenant-university-bioinformatics.

^{*}To whom correspondence should be addressed.

[†]The authors wish it to be known that, in their opinion, the first two authors should be regarded as Joint First Authors. Associate Editor: Zhiyong Lu

2 O.Falola et al.

Contact: ezekiel.adebiyi@covenantuniversity.edu.ng

1 Introduction

Genome-wide association studies (GWAS) analyses are widely used to report statistically significant genomic variants associated with a particular genetic trait or phenotype (Tam et al., 2019). However, in most cases, researchers are interested in understanding underlying molecular and biological functions that are triggered by these significant variants. Therefore, post-GWAS (pGWAS) analysis is required to address or interpret any GWAS findings (Gallagher and Chen-Plotkin, 2018). Though pGWAS analysis is critical for understanding the genetic mechanisms underlying many traits, it is challenging to perform pGWAS analysis for researchers with limited bioinformatics skills. These challenges are due to the complexity of installing some pGWAS tools, complex command line parameters, or the amount of Genomic data required for the pGWAS analysis. To facilitate pGWAS research, particularly for African bioinformatics and biomedical researchers due to the limited computing resources to store the huge annotated files and reference panels, we built SysBiol pGWAS as a web-based pGWAS tool.

2. SysBioPGWAS software

2.1. Input and data format

The input file for the SysBiolPGWAS pipeline is a GWAS summary file. A typical GWAS summary file contains nine fields which are: single-nucleotide polymorphisms (SNPs) ID, chromosome, genomic position, reference allele, alternative allele, beta score (effect size), standard error, z-score (summary statistics for SNP association with phenotype) and *P*-value. We used the standard GWAS summary files as described by Buniello *et al.* (2019) and MacArthur *et al.* (2021).

2.2 Individual tools and pipelines

SysBiolPGWAS provides direct access to several individual pGWAS tool pipelines. These tools are shown in Figure 1. The preprocessing pipeline step consists of cleaning up the input data; and utilizing the University of California, Santa Cruz (UCSC) LiftOver if needed (Haeussler et al., 2019). UCSC LiftOver is a major tool that is used for converting genomic coordinates between different assemblies. This tool is provided as a web-based tool hosted at the University of California, Santa Cruz (UCSC) Genome Browser (https://genome.ucsc.edu/cgi-bin/hgLiftOver). It is also available as a standalone tool (Luu et al., 2020).

The annotation pipeline combines four tools, which are Annovar tool (Wang et al., 2010), Ensembl Variant Effect Predictor (VEP) version 107 (McLaren et al., 2016), the Disease-specific Variant ANnotation tool (DIVAN) (Chen et al., 2016) and deTS software (Pei et al., 2019). The algorithm underlying these annotation tools is to determine what type of variant is being run and to assign an annotation score to the variants based on the source databases. Annovar tool is used to perform several functional scoring of variants. These functional scoring include the genome-wide annotation of variants that supports prioritization of non-coding variants by integrating various genomic and epigenomic annotations (GWAVA) (Ritchie et al., 2014), the functional prediction score generated by deep learning (DANN) (Quang et al., 2015), the functional prediction scores for mutations based on selective constraints across the human genome (GERP++) (Davydov et al., 2010; Cooper et al., 2005) and the spectral approach integrating functional genomic annotations for coding and non-coding variants (EIGEN) (Ionita-Laza et al., 2016). Also, Annovar tool is used for performing SNPs functional annotations and gene deleteriousness based on the dbNSFP database (Liu et al., 2020). On the other hand, Ensembl Variant Effect Predictor is used to perform functional prediction of variants using the Combined Annotation Dependent Depletion score (CADD) (Kircher et al., 2014; Rentzsch et al., 2019). DIVAN tool is

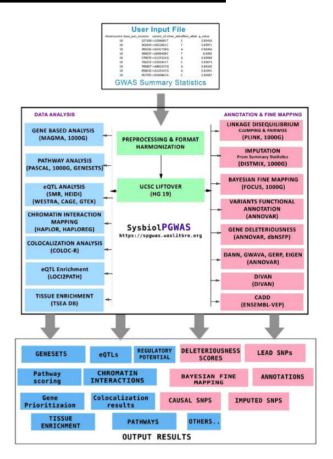


Fig. 1. SysBiolPGWAS pipelines and architecture, inputs and outputs. The rectangles in the center of the middle box show the preprocessing pipeline step consisting of cleaning up the input data and lifting over coordinates if needed. The rectangles to the right of the middle box indicate the group of tools that perform annotation of SNPs (detecting the type of SNPs) and fine-mapping (finding the causal SNPs) analysis. The rectangles to the left of the middle box show tools that perform omics based analysis

used for performing tissue-specific scoring for 45 different diseases/ traits. In addition, SysBiolPGWAS performs tissue-specific enrichment analysis for a list of genes that are associated with the variants using deTS (Pei *et al.*, 2019).

The estimating SNPs casualty pipeline step consists of finemapping and SNPs clumping. SysBiolPGWAS performs probabilistic fine-mapping, i.e. applied the Bayesian fine-mapping approach, using Fine-mapping Of CaUsal gene Sets software (Mancuso et al., 2019). Bayesian fine-mapping utilizes the statistical Bayes framework to estimate the probability of a given variant being a causal variant, i.e. estimating the Bayes factor (BF) (Wang and Huang, 2022). Such BF can be estimated from GWAS summary statistics such as variant P-value and its standard error of the effect even without accessing the individual-level genotype data (Mancuso et al., 2019). Approximation BF from GWAS summary report approximates Bayes factor (Wakefield, 2007, 2009). Maller et al. (2012) suggested estimating the probability of variant causality as a posterior inclusion probability (PIP) by using a simplified version of the Bayes model. Moreover, Maller et al. (2012) provided a method to estimate the smallest number of variants that can sum up to a predefined PIP threshold value and called it as the credible set. For detailed mathematical equations for the Bayesian fine mapping, refer to Wang and Huang (2022).

SysBiolPGWAS can select causal variants based on the linkage disequilibrium information in 1000 genomes using the clumping method of PLINK software. The process of variant clumping reports iteratively the most significant variant in the defined LD regions across the genome (Choi et al., 2020; Privé et al., 2019). In each LD region, the most significant variant, i.e. the SNP with the smallest P-value, is called the lead variant. However, the approach of choosing the lead variants is limited by the biological fact that the leads are not always considered as the causal variants (Schaid et al., 2018).

The step of omic-based analysis and reporting includes five types of analysis, which are (i) gene-level analysis, (ii) pathways analysis, (iii) eQTL analysis, (iv) chromatin interaction mapping [using Position Weight Matrices (PWM)] analysis and (v) colocalization analysis. SysBiolPGWAS uses MAGMA (Multi-marker Analysis of GenoMic Annotation) (de Leeuw et al., 2015) for performing genelevel analysis. The algorithm underlying MAGMA's gene level is based on the statistical regression approach and utilizing LD information to detect marker effects. SysBiolPGWAS pathways analysis using the Pathway Scoring Algorithm is implemented in Pascal software (Lamparter et al., 2016).

SysBiolPGWAS performs eQTL analysis by incorporating the GTEx multi-tissue eQTL information, which is very useful to understand the biological machinery underlying the variants in GWAS summary. The eQTL analysis can be performed using SMR and HEIDI (Wu et al., 2018), and Loci2Path tool (Xu et al., 2020). For the chromatin interaction mapping (via PWM) analysis, SysBiolPGWAS uses HaploR R package (Zhbannikov et al., 2017) to query HaploRegDB (Ward and Kellis, 2016) and RegulomeDB (Boyle et al., 2012). To perform colocalization analysis, SysBiolPGWAS uses coloc R package (Wallace, 2021).

The interpretation of several predicted variants in a GWAS to functional mechanisms is faced with several challenges identified by Gallagher and Chen-Plotkin (2018), a few of which include: (i) association of an SNP with a phenotype does not give sufficient information about the actual causal SNP or causal gene of that phenotype. (ii) Ninety percent (90%) of SNPs identified in a GWAS fall into a non-protein coding region (intergenic or intronic) which are very far from any known nearest gene. (iii) These SNPs that fall in noncoding regions may be enriched in putative regions of cis-regulatory elements (enhancers, silencers and promoters), however, because of the complex nature of regulation, it might be hard to associate these noncoding cis-regulatory elements (CREs) to correct target genes. We chose all these crop of tools in this study to help tackle these challenges. We also selected appropriate tools that help annotate variants in protein-coding and non-coding regions. In summary, each tool in the individual pipelines provides a unique pGWAS analysis to assist biologists in performing pGWAS analysis in one place, seamlessly overcoming the complex challenges and obtaining the results expeditiously.

2.3 Customized pipeline

This pipeline integrates several pGWAS tools and allows users to execute several of these tools in a single run. It should be noted that this pipeline performs pGWAS analysis only on the potential lead SNPs. First, we perform SNPs clumping to find lead SNPs, then we do gene-based analysis, followed by pathways analysis, eQTL analysis, SNPs annotations, deleteriousness and regulation analysis. We also execute Bayesian fine-mapping to report the probability of each SNP to be casual.

3 Conclusion

In conclusion, SysBiolPGWAS provides several pipelines consisting of multiple pGWAS tools with the current state-of-the-art annotation tools to perform complete pGWAS analysis and visualization for all users, especially scientists who are not command line or statistically inclined. This is the first version of the tool, and we plan to extend SysBiolPGWAS to include more comprehensive OMICS, further GWAS and pGWAS analysis by including more pGWAS

resources and tools. Online tutorials are available in the tutorial navigation link drop down section at https://spgwas.waslitbre.org/.

Acknowledgements

The authors acknowledge the editorial support of Mr. Babajide Ayodele at CUBRe.

Funding

Research reported in this publication was supported by the National Human Genome Research Institute (NHGRI), Office of the Director, National Institutes of Health (OD) under award numbers [U24HG006941 and U2RTW010679]. Furthermore, this work was supported by the World Bank funding for the ACE Impact projects. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health and the World Bank.

Conflict of Interest: none declared.

References

- Boyle, A.P. et al. (2012) Annotation of functional variation in personal genomes using RegulomeDB. Genome Res., 22, 1790–1797.
- Buniello,A. et al. (2019) The NHGRI-EBI GWAS catalog of published genome-wide association studies, targeted arrays and summary statistics 2019. Nucleic Acids Res., 47, D1005–D1012.
- Chen, L. et al. (2016) DIVAN: accurate identification of non-coding diseasespecific risk variants using multi-omics profiles. Genome Biol., 17, 252.
- Choi, S.W. et al. (2020) Tutorial: a guide to performing polygenic risk score analyses. Nat. Protoc., 15, 2759–2772.
- Cooper, G.M. et al.; NISC Comparative Sequencing Program. (2005) Distribution and intensity of constraint in mammalian genomic sequence. Genome Res., 15, 901–913.
- Davydov, E.V. et al. (2010) Identifying a high fraction of the human genome to be under selective constraint using GERP++. PLoS Comput. Biol., 6, e1001025.
- de Leeuw, C.A. et al. (2015) MAGMA: generalized gene-set analysis of GWAS data, PLoS Comput. Biol., 11, e1004219.
- Gallagher, M.D. and Chen-Plotkin, A.S. (2018) The post-GWAS era: from association to function. Am. J. Hum. Genet., 102, 717–730.
- Haeussler, M. et al. (2019) The UCSC genome browser database: 2019 update. Nucleic Acids Res., 47, D853–D858.
- Ionita-Laza, I. et al. (2016) A spectral approach integrating functional genomic annotations for coding and noncoding variants. Nat. Genet., 48, 214–220.
- Kircher, M. et al. (2014) A general framework for estimating the relative pathogenicity of human genetic variants. Nat. Genet., 46, 310–315.
- Lamparter, D. et al. (2016) Fast and rigorous computation of gene and pathway scores from SNP-Based summary statistics. PLoS Comput. Biol., 12, e1004714.
- Liu,X. et al. (2020) dbNSFP v4: a comprehensive database of transcript-specific functional predictions and annotations for human nonsynonymous and splice-site SNVs. Genome Med., 12, 103.
- Luu,P.L. et al. (2020) Benchmark study comparing liftover tools for genome conversion of epigenome sequencing data. NAR Genom. Bioinform., 2, lqaa054.
- MacArthur, J.A.L. et al. (2021) Workshop proceedings: GWAS summary statistics standards and sharing. Cell Genom., 1, 100004.
- Maller, J.B. et al.; Wellcome Trust Case Control Consortium. (2012) Bayesian refinement of association signals for 14 loci in 3 common diseases. Nat. Genet., 44, 1294–1301.
- Mancuso, N. et al. (2019) Probabilistic fine-mapping of transcriptome-wide association studies. Nat. Genet., 51, 675–682.
- McLaren, W. et al. (2016) The ensembl variant effect predictor. Genome Biol., 17, 122.
- Pei, G. et al. (2019) deTS: tissue-specific enrichment analysis to decode tissue specificity. Bioinformatics, 35, 3842–3845.
- Privé, F. et al. (2019) Making the most of clumping and thresholding for polygenic scores. Am. J. Hum. Genet., 105, 1213–1221.
- Quang, D. et al. (2015) DANN: a deep learning approach for annotating the pathogenicity of genetic variants. Bioinformatics, 31, 761–763.
- Rentzsch,P. et al. (2019) CADD: predicting the deleteriousness of variants throughout the human genome. Nucleic Acids Res., 47, D886–D894.

O.Falola et al.

- Ritchie, G.R. et al. (2014) Functional annotation of noncoding sequence variants. Nat. Methods, 11, 294–296.
- Schaid, D.J. et al. (2018) From genome-wide associations to candidate causal variants by statistical fine-mapping. Nat. Rev. Genet., 19, 491–504.
- Tam, V. et al. (2019) Benefits and limitations of genome-wide association studies. Nat. Rev. Genet., 20, 467–484.
- Xu,T. et al. (2020) Regulatory annotation of genomic intervals based on tissue-specific expression QTLs. Bioinformatics, 36, 690–697.
- Wakefield, J. (2007) A Bayesian measure of the probability of false discovery in genetic epidemiology studies. *Am. J. Hum. Genet.*, **81**, 208–227.
- Wakefield, J. (2009) Bayes factors for genome-wide association studies: comparison with P-values. *Genet. Epidemiol.*, 33, 79–86.
- Wallace, C. (2021) A more accurate method for colocalization analysis allowing for multiple causal variants. PLoS Genet., 17, e1009440.

- Wang,K. et al. (2010) ANNOVAR: functional annotation of genetic variants from high-throughput sequencing data. Nucleic Acids Res., 38, e164.
- Wang,Q.S. and Huang,H. (2022) Methods for statistical fine-mapping and their applications to auto-immune diseases. Semin. Immunopathol., 44, 101–113.
- Ward,L.D. and Kellis,M. (2016) HaploReg v4: systematic mining of putative causal variants, cell types, regulators and target genes for human complex traits and disease. *Nucleic Acids Res.*, 44, D877–D881.
- Wu,Y. et al. (2018) Integrative analysis of omics summary data reveals putative mechanisms underlying complex traits. Nat. Commun., 9, 918
- Zhbannikov,I.Y. et al. (2017) haploR: an R package for querying web-based annotation tools. F1000Res., 6, 97.