Linking genotype to phenotype in multi-omics data of small sample

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Abstract

Background: Genome-wide association studies (GWAS) that link genotype to phenotype represent an effective means to associate an individual genetic background with a disease or trait. However, single-omics data only provide limited information on biological mechanisms, and it is necessary to improve the accuracy for predicting the biological association between genotype and phenotype by integrating multi-omics data. Typically, gene expression data are integrated to analyze the effect of single nucleotide polymorphisms (SNPs) on phenotype. Such multi-omics data integration mainly follows two approaches: multi-staged analysis and meta-dimensional analysis, which respectively ignore intra-omics and inter-omics associations. Moreover, both approaches require omics data from a single sample set, and the large feature set of SNPs necessitates a large sample size for model establishment, but it is difficult to obtain multi-omics data from a single, large sample set.

Results: To address this problem, we propose a method of genotype-phenotype association based on multi-omics data from small samples. The workflow of this method includes clustering genes using a protein-protein interaction network and gene expression data, screening gene clusters with group lasso, obtaining SNP clusters corresponding to the selected gene clusters through expression quantitative trait locus data, integrating SNP clusters and corresponding gene clusters and phenotypes into three-layer network blocks, analyzing and predicting based on each block, and obtaining the final prediction by taking the average.

Conclusions: We compare this method to others using two datasets and find that our method shows better results in both cases. Our method can effectively solve the prediction problem in multi-omics data of small sample, and provide valuable resources for further studies on the fusion of more omics data.

Keywords: Multi-omics, Small sample, SNP, Gene, Phenotype

Background
An important goal of current genetics is to establish global functional associations between genotype and phenotype, the so-called genotype-phenotype map [1]. The study of genotype-phenotype association sheds light on the process of genetic variation [2]. Genome-wide association studies (GWAS) that link genotype to phenotype represent an effective way to associate individual genetic backgrounds with specific diseases or traits. The strategy is to locate all the differential sites in a genome and correlate them to a phenotype. Over the past decade, many GWAS have been performed to identify genetic variants associated with complex human diseases or traits. These findings highlight novel associations between variants and traits, provide insight into racial variations in complex traits, and lead to multiple clinical applications [3, 4]. However, most variants explain only a small fraction of the causal genetic factors. According to the principle of GWAS, although thousands of single nucleotide polymorphisms (SNPs) for complex diseases and traits have been identified, single-omics analysis can only provide limited information on biological

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mechanisms, and the functions and mechanisms of SNP loci remain largely unclear.

Due to limitations at the single-omics level, it is necessary to integrate multi-omics data to more accurately predict the biological associations between genotype and phenotype [5]. Such data allow us to study interactions across omics, and provide opportunities to further examine genotype-phenotype associations and uncover the underlying mechanisms [6]. Gene expression data are typically incorporated to analyze the impact of SNPs on phenotype [7–9]. The two main approaches of multi-omics data integration are multi-staged analysis and meta-dimensional analysis [10, 11].

When using a multi-omics biological network to explore genotype-phenotype association, it is generally believed that the difference in phenotypic traits is primarily attributable to the cumulative effect of omics. For example, SNPs lead to changes in gene expression, which in turn affect protein expression and ultimately cause diseases [12]. Such layer-by-layer integration analysis is usually called multi-staged analysis. The idea is to establish the association between two layers by linear regression, partial least squares (PLS) [13, 14], canonical correlation analysis, and correlation coefficients, and to predict diseases through the associations across layers. Multi-staged analysis bridges the gap between genotype and phenotype by gene expression traits. The most commonly used such approaches are three-layer methods such as the network-driven association mapping (NETA M) algorithm for regression models based on biological networks [15], in which a three-level association network is constructed, linear regression is used to establish the association between SNPs and genes, and logistic regression is used to link genes with phenotypes (with values 0 or 1, representing the absence or presence, respectively, of a disease). Disease prediction by analyzing the effect of SNPs on gene expression is more accurate than direct prediction by SNPs. This confirms that the three-layer network is more likely to reflect real biological relationships. Similar models include PrediXcan [16] and backward three-way association mapping (BTAM) [17]. However, when a regression model is established on a three-layer network, intra-omics correlations are not considered, which results in low model accuracy.

Meta-dimensional analysis has the three approaches of concatenation-based, transformation-based, and single-omics model-based integration. In concatenation-based integration, the features of omics data are integrated through machine learning to form a more comprehensive input matrix, through which a prediction model is established [18]. In transformation-based integration, multi-omics data are transformed to an intermediate form, which is used for integration and development of a predictive model [19]. In single-omics model-based integration, predictive models are established using individual omics data, and their predictions are integrated to produce the final output [20]. A common drawback of meta-dimensional analysis is that the omics data have identical weights, meaning that they are used to predict phenotypes from different perspectives without examining their biological correlations. Although meta-dimensional analysis can improve the accuracy of phenotype prediction, the biological significance of multi-omics data integration remains unclear.

Furthermore, genotype-phenotype association by multi-omics requires omics data from a single sample set, and because of the large feature set of SNPs, both multi-omics analysis methods require larger sample sizes for model construction. Acquisition of clinical data can be hampered by patient privacy protection and specific data requirements of various institutions. Therefore, public clinical data do not meet the needs of multi-omics data integration methods in terms of sample size and number of omics. We propose a method of genotype-phenotype association for multi-omics data based on a small sample size and large feature set. This method offers the following innovations. a) It solves the problem of ineffective regression due to a large feature set and small sample size in three-layer networks. b) It considers intra-omics associations to improve prediction accuracy. c) It analyzes the biological pathway associations across omics layers to clarify the biological significance. d) It accounts for tissue specificity, because each tissue of a multicellular organism has distinguishing characteristics due to the tissue-specific expression of genes that confer unique morphological structures and physiological functions on various tissues [21, 22]. e) In the study of the SNP-gene-phenotype pathway, our method is the most time-saving over state-of-the-art methods.

Results
Data sources and preprocessing
Two datasets derived from the Gene Expression Omnibus (GEO) database were used to verify the effectiveness of our method [23]. GSE33356 represents the data of lung adenocarcinoma [24]. Affymetrix SNP 6.0 and Affymetrix U133 Plus 2.0 microarrays were used to analyze the specimens of lung tumors and normal tissues from 84 nonsmoking women with adenocarcinoma. GSE114269 represents the data comparing medullary breast carcinoma (MBC) with non-medullary basal-like breast carcinoma (non-MBC BLC) among 48 patients [25]. These two datasets were selected to show whether our method can be useful in genotype and phenotype classification in various cases of small sample sizes and large feature sets.
The PPI network data were derived from PICKLE (Protein InterAction KnowLedgебase) [26], which are metadata that integrate human PPI from various open sources through gene ontology information. The eQTL data were obtained from GTEx Analysis V7 (dbGaP Accession phs000424.v7.p2) [27]. For accurate data prediction, eQTL data were selected based on tissue specificity. For example, lung eQTL data was selected for the first dataset, and breast tissue eQTL data for the second.

The data were preprocessed for consistent nomenclature of SNPs and genes in data of various types. Data with more than 10% missing values were removed from SNPs, data with less than 10% missing values were imputed with the most frequent value or the mean value, and only SNP data with minor allele frequency (MAF) greater than 0.1 were used.

**Prediction analysis**

There is currently no typical method of genotype-phenotype association for data with a small sample size and a large number of features. We compared the following methods to verify the feasibility of our approach. In comparison experiments, the method developed in this paper was named GSPLS (Group lasso and SPLS model). The detailed implementation process of this method can be viewed in the section “Methods” and the Supplementary Information.

1) The method GGLM (Group lasso and Generalized Linear Model) differs from GSPLS in that multiple regression rather than SPLS is used to associate SNP layers with gene layers, which allows us to verify the effect of gene-gene association while considering inter-omics association.

2) The method NETAM [15] involves multi-staged analysis, in which gene clustering are absent. Each SNP group was defined by the SNPs within the transcribed region of a gene, and a large three-layer network is established based on all the data and subjected to direct multiple regression with lasso. This method does not cluster SNPs and genes, and can be used to verify the effect of inter-omics association without gene associating and clustering. NETAM consists of two parts: Sparse Regression and Stability Selection. There are two user-defined parameters $T$ and $\pi_{thr}$ in this method ($\pi_{thr}$: threshold for stability selection, $T$: total number of random samples). These two parameters serve the Stability Selection part. However, due to our small sample size, Stability Selection cannot be carried out, so we test NETAM without the benefits of stability selection, where lasso and L1-regularized logistic regression are employed with 5-fold cross-validation.

3) The method mixOmics [28] involves multi-omics integration, in which SNP and gene data are used to independently establish prediction models, and predictions are integrated to generate the final output. This method ignores SNP-gene association and allows us to verify the effect of omitting inter-omics association, but it fails to analyze the inter-omics pathway relationship.

Our method was compared to the above three approaches on GSE33356 and GSE114269 datasets. Because it was a dichotomous classification, receiver operating characteristic curves were used for comparison and explanation. Figure 1 shows the area under curve for the four methods.

As shown in Fig. 1, our method achieved better results on both datasets. The performance of GSPLS was significantly improved over that of GGLM, indicating that inter-omics associations should be considered in the analysis to better fit real biological systems. Both GSPLS and GGLM with gene network clustering showed superiority over other methods, indicating that clustering and screening gene data followed by taking the average are suitable for processing data from small samples. The NETAM method showed the worst performance, possibly because it is not suitable for small samples, but it provides guidelines for inter-omics pathway analysis. It has been documented that when analyzing SNP-gene-phenotype associations, this algorithm is suitable when the sample size reaches 500. In the case of a small sample, it is ineffective in regression. The result of mixOmics was similar to that of GGLM, especially on the second dataset. In our test, it sometimes even outperformed GGLM, but this method ignores inter-omics associations.

In this experiment, we found that the results of the last three methods were unstable with small sample sizes. Overall, GSPLS was superior to GGLM, mixOmics, and NETAM with small sample sizes.

Through the selection of gene clustering and corresponding SNP clusters, we not only grouped the sample features, but also screened the grouped features through Group Lasso, so that the feature quantity contained in each block was sharply reduced, which not only reduced the sample size required by our algorithm, but also reduced the computation time. We tested the four methods GSPLS, GGLM, NETAM and mixOmics on GSE33356 data with 10 repeated experiments respectively on a PC with Intel Core i7-10510U processors, 16 Gb RAM, and NVIDIA GeForce MX330 video card. The computation time of GSPLS was 66.213 ± 3.905 s, GGLM was 138.527 ± 12.149 s, NETAM was 712.141 ± 89.491 s and mixOmics was 47.065 ± 4.135 s. Although the GSPLS time is longer than the mixOmics time, the mixOmics cannot reflect the pathway correlation. In the study of the SNP-gene-phenotype pathway, our method is the most time-saving.

**Sample sizes analysis**

In the study of three-way association analysis among genotypes, gene traits and phenotypes, large number of...
samples are often needed. For example, NETAM [15] showed significantly better performance (larger area under the curve) for $N > 200$, but GSPLS only needs dozens of samples.

To validate the effect of sample size, we randomly select samples $N = 80, 60, 40, 30, 20$, from the data set GSE33356, and $N = 48, 40, 30, 20$, from the data set GSE114269 (Because this data set contains only 48 sample sizes). The positive and negative sample sizes are guaranteed to be equal during extraction. When sample size is less than 20, Group Lasso could not achieve effective regression for these two data sets. Therefore, the minimum sample size we choose is 20. In Fig. 2, we show receiver operating characteristic (ROC) curves that show true positive and false positive rates of the results produced by GSPLS with different sample sizes. The results show that the general trend of AUC value decreases with the decrease of sample size. Although this

Fig. 1 ROC curves to compare the performance of GSPLS with association mapping methods such as GGLM, NETAM, and mixOmics. The left one is the result of analysis on GSE33356, and the right one is the result of analysis on GSE114269. The legend of each figure contains the AUC value of each method on the data. The AUC value of our method is the highest, indicating that our method is superior to other methods.

Fig. 2 ROC curves to compare the performance of GSPLS with different sample sizes. The left one is the result of analysis on GSE33356, and the right one is the result of analysis on GSE114269. The legend of each figure contains the sample sizes and the AUC value of each sample size on the data. With the decrease of the sample size, the corresponding AUC value also showed a decreasing trend.
method is suitable for small sample data, unlike NETA
M, which requires a sample size of more than 200, it
also requires a sample size of more than 20.

**Pathway analysis**
We analyzed the generated pathways to further demon-
strate the effectiveness of our method. The weight of the
edge between each gene and relevant SNP in the cluster
was obtained by SPLS, and the weight of the edge con-
necting disease and the relevant gene was obtained by
group lasso. The two layers of weights were multiplied
to yield the weight value of a pathway (if no edge was
present, then the edge weight was set to zero), which we
name Pathway Score (PS) to reflect its importance.

Using GSE33356 as an example, we ranked the gener-
ated associations by PS, and aligned the top 50 pathways
with the lung cancer-associated SNPs and genes in the
PhenoScanner database [29], which contains 137
genotype-phenotype association datasets, as well as asso-
ciation catalogs such as NHGRI-EBI GWAS, NHLBI
GRASP, and dbGaP. Alignment against PhenoScanner reflects the distribution of SNPs or genes in the path-
ways obtained by this method (Fig. 3).

As shown in Fig. 3 and Table 1, We found that the
percentage of SNPs or genes present in PhenoScanner
decreased as the total number of PS ranks increased,
mainly because when numerically solving pathway rela-
tionships, SNPs or genes with the same trend are ran-
domly interchanged during data sparsification. With the
decrease of PS, the total number of SNPs and genes with
the same trend increased, the possibility of interchanging
SNPs or genes matching with the database increased,
and thus the percentage of such SNPs or genes
decreased.

**Discussion**
We addressed genotype–phenotype association based on
multi-omics data for small samples. In the case of data
with numerous features, there are two commonly used
approaches: dimension reduction and feature selection.
However, the former may lead to the inability to analyze
features before dimension reduction. Therefore, sparsifi-
cation and existing associations were used to screen and
reduce the number of features. The greatest contribution
of this paper is the ability to group genes into clusters
according to their network characteristics while signifi-
cantly reducing the number of features in each cluster.
The clustering and effective screening of gene networks
drastically reduces the number of features of the three-
layer network, making it suitable for application to data
with small sample sizes.

Data is a key in multi-omics studies. However, the fol-
lowing two conditions generally occur to the multi-
omics data. One is that with the increase of the types of
omics data, the multi-omics data is not easy to obtain
and the data sample size is not large. In this paper, such
problems can be solved in the case of small samples, the
advantage is that it can reflect the association relation-
ship of each omics, it makes the biological mechanism
clearer, but the accuracy is not high. The other is that
when the sample size is large, the number of omics types
is often small, such as GWAS data. Although GWAS
analysis has a certain accuracy, it is generally analyzed
by statistical methods without understanding the in-
ternal biological mechanism. How to achieve the analysis
effect of multi-omics data, when we use the data with
sufficient sample size and insufficient omics types. This
will be the direction of our next research.

Our method has obvious advantages, but the data rely
on the PPI network and eQTL association. Therefore,
only associations in an SNP/CNV-gene-phenotype
three-layer network can be processed, rather than any
three omics, such as SNP, methylation, and phenotype.

When clustering with SPICi, although we can select
the ranges of the three clustering hyperparameters ac-
cording to certain criteria, unknown factors still
influence the selection of range limits, and this method is highly sensitive to the choice of hyperparameters. We also attempted to replace SPICi clustering with fuzzy clustering, which showed results similar to sampling multiple gene clusters with replacement and was expected to achieve better clustering. However, fuzzy clustering requires specification of parameters, including cluster number. Inappropriate initial parameters may affect clustering accuracy, and the real-time performance of the algorithm can be compromised by a large sample dataset and feature set. These problems must be addressed in future studies.

**Conclusion**

In summary, in this study we used SPICi and Group Lasso methods to perform gene cluster and screen, obtained the SNP corresponding to gene in each cluster through eQTL data, established the association relationship between SNP and gene by SPLS, and formed block together with phenotype. Finally, these blocks are integrated by taking the average. Compare with state-of-the-art methods, our method achieved better results on GSE33356 and GSE114269 datasets, and it consumes the least time. To validate the effect of sample size, we randomly select samples from these two datasets, we found that our method was effective for the sample size greater than 20. And we ranked the generated associations by PS, SNPs or genes in 6 of the top 10 pathways were present in PhenoScanner, as were those in 10 of the top 20 pathways. Collectively, our method can effectively solve the prediction problem in multi-omics data of small sample, and provide valuable resources for further studies on the fusion of more omics data.

**Methods**

**Algorithms**

Multi-staged analysis ignores intra-omics correlations and requires a large sample size, whereas meta-dimensional analysis ignores inter-omics associations. To address this issue, we propose an approach to genotype-phenotype association for small samples based on multi-omics data. The model is illustrated in Fig. 4. Diseases are not caused by the effects of one or more genes [30–32], but by gene interactions. A protein–protein interaction (PPI) network can reflect gene–gene associations, but this network is unweighted. Gene expression data can be used to calculate the Pearson correlation coefficient between genes, which serves as the weight of an edge, and this can be combined with a PPI network to produce a weighted gene network. Closely related genes are more likely to interact and impact on a phenotype, so we performed speed and performance in clustering (SPICi) on a gene association network to obtain multiple gene clusters (see section “Clustering method selection and hyperparameter setting”) [33]. Due to the large number of genes, and thus the many obtained gene clusters, group lasso regression can be performed on gene clusters and phenotypes. Gene clusters with nonzero coefficients are most likely to have impacts on a disease, and this regression serves the purpose of gene cluster screening. SNP clusters corresponding to the selected gene clusters can be identified by expression quantitative trait loci (eQTL) that control the expression level of quantitative trait genes. It is important to note that the same SNP may occur in multiple SNP clusters. In this way, correlated SNP clusters, gene clusters, and phenotypes are combined in a three-layer network, which is called a block. The SNP and gene layers can be linked by sparse partial least squares (SPLS) [14], and gene layers and phenotypes by logistic regression. The prediction and analysis of blocks are integrated by taking the average. The flowchart is shown in Fig. 5. Additional details are provided in the Supplementary Information.

**Clustering method selection and hyperparameter setting**

Clustering algorithms play an important role in the analysis of biological networks and can reveal their functional modules. Most clustering algorithms perform well...
on biological networks of moderate size, but they become impractical on larger networks due to low speed or high network complexity. According to the analysis of Jiang et al. for each clustering methods including Markov cluster (MCL) [34], molecular complex detection (MCODE) [35], Cfinder [36], the memory-constrained unweighted pair group method using arithmetic averages (MCUPGMA) [37], and dense module enumeration (DME) [38], SPICi [33] was selected to cluster the generated gene network. SPICi uses a heuristic approach to greedily build clusters, which is orders of magnitude faster, and was the only method to successfully cluster all test networks in a short period of time [33]. Its three hyperparameters are minimum cluster size, minimum support threshold, and minimum cluster density, which jointly affect the number of clusters and the number of elements in each cluster. To facilitate subsequent analyses, the final numbers of clusters and elements should fall in appropriate ranges. If there are too many elements in a cluster, during group lasso calculation the goal-
oriented error caused by a given penalty parameter is greater for a group with more elements. If there are too few elements in a cluster, the effect of gene association on disease cannot be effectively analyzed. Generally, gene data were clustered into 100–200 clusters, and each cluster containing between 5 and 100 genes. We further analyzed the settings of the three hyperparameters as they have different characteristics.

The minimum cluster size is used to determine the inclusion/exclusion of a cluster by comparing the number of genes involved in it. If the number of elements in a cluster is greater than the minimum cluster size, then the cluster is included. If the minimum cluster size is too small, then it fails to capture the associations between genes, and the opposite may result in the inappropriate deletion of clusters. According to tests on different datasets, the minimum cluster size was set to the interval of [13, 14].

In an undirected graph $G = (V, E)$, for any vertex $u$ and connecting vertex set $S \subset V$, support is defined as

$$support(u, S) = \sum_{v \in S} w_{u,v},$$

i.e., the sum of weights of all edges connected to vertex $u$, where $w_{u,v}$ is the weight of any edge $(u, v) \in E$. If the support of a vertex is less than the minimum support threshold, then the vertex is discarded. The weight of an edge is represented by the Pearson correlation coefficient of two vertex vectors; a coefficient can be positive or negative, and coefficients can cancel each other during the calculation of support. Therefore, the absolute value of the Pearson correlation coefficient was used, so $w_{u,v} \in (0, 1]$. Generally, the absolute value of Pearson correlation coefficient is less than 0.2, which indicates very weak correlation or no linear correlation, so we only take the edges whose absolute value of Pearson correlation coefficient is greater than 0.2 to form the gene-gene network. If a vertex is only linked to a gene with weak association, then additional noise may be introduced, so it is required that if a vertex is only linked to genes with weak association, then two or more such edges should be present, so the lower limit of the minimum support threshold was set to 0.4. In tests with various datasets, when the minimum support threshold value was greater than 0.7, the total number of genes became insufficient to reflect the impact of gene

|                  | minimum cluster density | minimum support threshold | minimum cluster size |
|------------------|-------------------------|---------------------------|----------------------|
| GSE33356         | 0.1                     | 0.4                       | 5                    |
| GSE114269        | 0.2                     | 0.5                       | 5                    |

Fig. 6 Model of genotype-phenotype association based on biological network. The left one is the schematic diagram without considering the intra-omics associations, and the right one is the schematic diagram used in our method. It contains both intra-omics associations and inter-omics associations, which can better reflect the biological reality. Without considering the intra-omics associations, the model cannot truly reflect the biological reality, and without considering the inter-omics associations, it cannot well reflect the whole biological system.
association on disease. Therefore, the minimum support threshold was set to the interval of [0.4, 0.7].

Cluster density is defined as the sum of edge weights divided by the total number of possible edges, and it is used to reflect the density of subgraphs. It is calculated as

$$density(S) = \frac{\sum_{u,v \in S} W_{u,v}}{|S| \times (|S| - 1)/2}.$$  

As shown in the formula, too small a density increases the number of elements in each cluster and decreases the total number of clusters. When the density of a cluster is less than the minimum cluster density, SPICluster it into two or more clusters. Therefore, the minimum cluster density directly affects the total number of clusters and has the greatest impact on clustering among the three hyperparameters. Based on comparative experiments, the minimum cluster density was set to [0.1, 0.6], and the parameter was tested in increments of 0.1 during the experiment. Through testing, we determined the hyperparametrical settings of the two data sets in our paper, as shown in Table 2.

Three-layer network construction

Gene clusters closely related to a disease can be selected by group lasso [39, 40]. The causal link from these gene clusters to a disease is established by related differential sites in the genome. The corresponding SNP clusters can be obtained by eQTL data. Each SNP cluster as well as the correlated gene cluster and phenotype can be combined in a three-layer network block. After dividing into blocks, the number of SNPs and gene features in the three-layer structure decreases drastically, as does the sample size required for effective regression, which facilitates regression of numerous features with small sample size. When analyzing the three-layer structure of each block, although interlayer regression alone can predict some pathway relationships, it does not consider intra-omics associations, and it deviates from biological reality. However, to only consider intra-omics associations and ignore inter-omics pathway correlations fails to reflect the whole biological system and represents only the local situation (Fig. 6). Therefore, when we look at the associations between SNPs and genes, the associations of individual elements between layers (left part of Fig. 6) were expanded to intra- and inter-layer associations (right part of Fig. 6). That is, the many-to-one associations were broadened to many-to-many associations. Accordingly, the solution was altered from multiple regression to SPLS, which incorporates a penalty function in PLS [14]. The principle of PLS is as follows. With $q$ dependent variables $\{y_1, ..., y_q\}$ and $p$ independent variables $\{x_1, ..., x_p\}$ [41], the statistical relationships between dependent and independent variables can be studied by observing $n$ sample points, which constitute the data matrices $X(n \times p)$ and $Y(n \times q)$ of independent and dependent variables, respectively. PLS regression extracts principal components $t_1$ and $u_1$ from $X$ and $Y$, respectively. For regression analysis, $t_1$ and $u_1$ should carry as much information as possible of the data they represent. After extraction of the first set of principal components $t_1$ and $u_1$, PLS performs regression of $X$ on $t_1$ as well as $Y$ on $u_1$. If the regression equations are sufficiently accurate, then the algorithm is terminated. Otherwise, a second round of component extraction is performed using the residual information after removing the variance of $X$ and $Y$ explained by $t_1$ and $u_1$, respectively. This process is repeated until satisfactory accuracy is achieved. SPLS combines the advantages of principal component analysis, canonical correlation, and linear regression, and effectively solves the problems of ineffective regression and multicollinearity among features due to fewer samples than features. In order to assess the impact and usefulness of the single steps, GSE33356 is used as an example data to analyze the intermediate results of each step in the Supplementary Information.

Abbreviations

GWAS: Genome-wide association studies; SNPs: single nucleotide polymorphisms; BTAM: backward three-way association mapping; eQTL: expression quantitative trait loci; SPLS: sparse partial least squares; GEO: Gene Expression Omnibus; MBC: medullary breast carcinoma; non-MBC: non-medullary breast carcinoma; PICKLE: Protein InteraCtion KnowLedgebasE; MAF: minor allele frequency; ROC: receiver operating characteristic; PS: Pathway Score

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s12864-021-07867-w.

Additional file 1.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (No. 61772426). We thank the Data mining and Bioinformatics Lab of NWPU for technical support.

Authors’ contributions

XG and XS conceived the research; XG, YS and SL developed the method; XG and XS performed the data analyses; XG and YQ wrote or edited the manuscript. All authors read and approved the final manuscript.

Funding

The publication costs for this article were funded by Air Force Engineering University.

Availability of data and materials

The method is available at https://github.com/2017100647/GSPLS. Publicly available datasets were analyzed in this study. This data can be found here: https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE33356; https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE114269; http://www.pickle.gr/; https://gtexportal.org/home/datasets.
Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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Received: 24 February 2021 Accepted: 30 June 2021

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