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Importance-Penalized Joint Graphical Lasso (IPJGL): differential network inference via GGMs

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Abstract

Motivation: Differential network inference is a fundamental and challenging problem to reveal gene interactions and regulation relationships under different conditions. Many algorithms have been developed for this problem; however, they do not consider the differences between the importance of genes, which may not fit the real-world situation. Different genes have different mutation probabilities, and the vital genes associated with basic life activities have less fault tolerance to mutation. Equally treating all genes may bias the results of differential network inference. Thus, it is necessary to consider the importance of genes in the models of differential network inference.

Results: Based on the Gaussian graphical model with adaptive gene importance regularization, we develop a novel Importance-Penalized Joint Graphical Lasso method (IPJGL) for differential network inference. The presented method is validated by the simulation experiments as well as the real datasets. Furthermore, to precisely evaluate the results of differential network inference, we propose a new metric named APC2 for the differential levels of gene pairs. We apply IPJGL to analyze the TCGA colorectal and breast cancer datasets and find some candidate cancer genes with significant survival analysis results, including SOST for colorectal cancer and RBBP8 for breast cancer. We also conduct further analysis based on the interactions in the Reactome database and confirm the utility of our method.

Availability and implementation: R source code of Importance-Penalized Joint Graphical Lasso is freely available at https://github.com/Wu-Lab/IPJGL.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

As biological data becomes more accessible, researchers are looking to extract key information from the differences in data between normal and disease samples. Although many studies analyze differences between data and extract knowledge from a single-gene perspective (Love et al., 2014; Robinson et al., 2010), there is still a wealth of information to be mined in the differential data. Studies have shown that the co-expression pattern between a pair of genes may have changed significantly even in the absence of significant differences in the expression levels of either one of these genes (de la Fuente, 2010). Therefore, it is necessary to study biological mechanism from the perspective of gene interactions. And studies have shown that biological networks differ a lot in different states, different organizations, and even at different times (Bandyopadhyay et al., 2010; Greene et al., 2015; Ha et al., 2015). Therefore, in order to study key genes which are the root cause of diseases, we need to focus on the differential networks. In general, we focus more on the differential expression between normal samples and disease samples, i.e. given two groups of gene expression data, our problem becomes how to build a differential co-expression network to explore the key genes by differential network analysis (DiNA). However, it is not simple, because there are always numerous false positives in the network inferred by various methods. One of the reasons is that many interactions in co-expression networks result from indirect influence (Cecchini et al., 2018). To eliminate such indirect influence, many models have been developed. One kind of the models widely developed by researchers is the Gaussian graphical models (GGMs) because of its good property: if the entry of the precision matrix Θij is zero, it is equivalent that variables i and j are conditionally independent, i.e. there is no co-expression interactions between i and j (Friedman et al., 2008). Many researchers have conducted research based on the GGMs and proposed various methods to meet different assumptions.

Under usual circumstances, the number of genes p ≥ n, the number of samples, which results in the sample covariance matrix not
being invertible. This means the precision matrix $\Theta = \Sigma^{-1}$ could not be computed directly, where $\Sigma$ is the real covariance matrix. Graphical Lasso (GLasso) was a popular method to estimate the precision matrix $\Theta$ (Banerjee et al., 2008; Friedman et al., 2008; Yuan and Lin, 2007). This method added the Lasso penalty term on the likelihood function to obtain a sparse solution, guiding the development of the GGMs. Subsequently, many researchers have researched the differential networks based on this model. Danaher et al. (2014) first used the differential matrix as the penalty term assuming that the networks from two groups should be similar in the overall structure despite the relatively subtle differences and hence proposed Joint Graphical Lasso (JGL). On this basis, Mohan et al. (2014) proposed Perturbed-Node Joint Graphical Lasso (PNJGL) and Co-hub Node Joint Graphical Lasso (CNJGL) to solve node-based problems: the first one assumes that there are hub nodes in differential networks, and the second assumes that different states have the same hub nodes. The differential network can not only be obtained by the direct subtraction between networks of two states but can also be directly solved as a variable (He and Deng, 2019; Tang et al., 2020; Tian et al., 2016; Yuan et al., 2017; Zhao et al., 2014), on the premise of ensuring accuracy while greatly reducing the number of samples required.

However, the above methods have a common shortcoming, i.e. they put all genes in the same position. This is not the case in the real world. Different genes have different mutation probabilities. Even the same gene has different mutation rates in different tissues and locations (Scally, 2016). In general, the more important genes are less likely to mutate. This is because, on the one hand, if extremely important genes are mutated in an individual, such as genes that regulate essential activities of life, the individual will have a high probability of not being sampled due to rapid death. On the other hand, studies have shown that even if a mutation occurs, important genes will be repaired preferentially, i.e. one of the reasons for the different probabilities of gene mutations is the DNA repair mechanism (Supek and Lehner, 2015). Some methods adjust the weights of different genes (Lyu et al., 2018; Ou-Yang et al., 2020; Zuo et al., 2017). However, these methods need the prior information from the database. Once the prior information is insufficient, or the information in the database is a little biased, the results may also have certain deviations. Some method does not rely on prior information (Salamangov et al., 2019), but it requires the time-consuming iterative generation of weights; moreover, it only considers a single state, which is not developed for inferring differential networks.

To address the above issues, we proposed a novel differential network inference method to meet the hypothesis that the important genes are less likely to mutate. In this proposed model, we add an importance penalty to each gene, and the penalties vary with the solution. The genes that are significantly altered between two states will be penalized if they also have large degrees in either one of two estimated networks. The comprehensive simulation experiments proved that the proposed Importance-Penalized Joint Graphical Lasso (IPJGL) method outperforms several state-of-the-art differential network inference methods based on GGMs, on the datasets simulated under realistic and reasonable assumptions. To better evaluate the results of differential network inference, we also propose a new metric named APC2 for the differential levels of gene pairs. We further applied IPJGL to the TCGA colorectal cancer and breast cancer datasets. The networks obtained by IPJGL showed some connections within the same family, as well as those already existed in BioGRID (Stark et al., 2006) and Cytoscape Reactome database (Jassal et al., 2020; Shannon et al., 2003; Sidiropoulos et al., 2017; Wu et al., 2010), which confirmed the validity of our method. Moreover, in the enrichment analysis, our method successfully obtained a gene set among about 1000 genes, which was significantly enriched in the pathway most relevant to the disease. On a deeper level, through survival analysis, we discovered a gene SOST which may be closely related to colorectal cancer. It is an inhibitor of WNT signaling (Wang et al., 2016), and WNT signaling happens to have a signaling pathway related to colorectal cancer (Clevers, 2006). Research on these genes may reveal the pathological mechanism of colorectal cancer.

## 2 Materials and methods

### 2.1 JGL and its variants

Given the gene expression matrix of two sets of samples, normal samples, $X^{(1)} = (x_{ij}^{(1)})_{n_1 \times p}$, and disease samples, $X^{(2)} = (x_{ij}^{(2)})_{n_2 \times p}$, where $p$ represents the number of genes, and $n_1$ and $n_2$ represent the number of normal samples and disease samples, respectively. Let $S^{(1)} = (s_{ij}^{(1)})_{p \times p}$ and $S^{(2)} = (s_{ij}^{(2)})_{p \times p}$ be the corresponding sample covariance matrices, $\Theta^{(1)} = (\theta_{ij}^{(1)})_{p \times p}$ and $\Theta^{(2)} = (\theta_{ij}^{(2)})_{p \times p}$ be the corresponding precision matrices. Also, we define the matrix norms, $\|X\|_1 = \sum_{i,j=1}^p |x_{ij}|$ and $\|X\|_2 = \sum_{i=1}^p x_{ii}^2$. For vectors $V = (v_1, v_2, \ldots, v_p)$, the vector norm is defined as $|V|_1 = \sum_{i=1}^p |v_i|$ and $|V|_2 = \sqrt{\sum_{i=1}^p v_i^2}$.

Under the assumption that the variable $X = (x_1, x_2, \ldots, x_p)$ are from the $p$-dimensional Gaussian distribution, Danaher et al. (2014) proposed the JGL model. They followed Friedman et al. (2008) and added a penalty term to make the precision matrices of the two states share an overall similar structure as follows:

\[
\min_{\Theta^{(1)}, \Theta^{(2)}} -L(\Theta^{(1)}, \Theta^{(2)}) + \frac{1}{2} \sum_{k=1}^2 \sum_{i,j} s_{ij}^{(k)} \theta_{ij}^{(k)} + \frac{1}{2} \lambda_2 \|\Theta^{(1)} - \Theta^{(2)}\|_F^2,
\]

where $\Theta^{(k)}$ is the space of $p$-dimensional positive definite matrices and $L(\Theta^{(1)}, \Theta^{(2)}) = \sum_{i=1}^p n_i \log(\det(\Theta^{(i)})) - tr(S^{(i)}(\Theta^{(i)}))$ is the likelihood function of the GGMs. The first penalty term is the sparsity penalty. The second penalty term is the similarity penalty, which is defined as

\[
P(\Theta^{(1)}, \Theta^{(2)}) = \sum_{i=1}^p |\theta_{ii}^{(1)} - \theta_{ii}^{(2)}|.
\]

This penalty term penalizes edges that are different between two states, and its weight is $\lambda_2$.

On this basis, Mohan et al. (2014) proposed PNJGL and CNJGL, of which the first term is the same likelihood function, but the first penalty is slightly different in that they penalized the diagonal values, i.e. $\sum_{i=1}^p \sum_{j=1}^p |\theta_{ij}^{(i)}|$, and the second penalty is modified as

\[
P(\Theta^{(1)}, \Theta^{(2)}) = \sum_{i=1}^p |\theta_{ii}^{(1)} - \theta_{ii}^{(2)}|\cdot
\]

and

\[
P(\Theta^{(1)}, \Theta^{(2)}) = \sum_{i=1}^p |\theta_{ii}^{(1)} - \theta_{ii}^{(2)}|\cdot
\]

where $q = 1.2$, respectively. They decomposed the matrix symmetrically to ensure that there are hub nodes in the corresponding network structure. PNJGL makes the hub structures exist in the differential matrix, and CNJGL supposes two states have the same hub nodes. In these models, all genes are treated equally, which does not tally with the actual situation.

### 2.2 Importance-Penalized Joint Graphical Lasso

We proposed the IPJGL model based on the following assumptions: (i) the expressions of each gene in the samples of each state follow a Gaussian distribution; (ii) the importance of genes is correlated with their degrees in the network and the important genes are less likely
to mutate; (iii) if a gene mutates, its existed or new interactions with other genes are dropped or established with certain probabilities, respectively.

The IPJGL model is given as follows:

$$\min_{\Theta^{(1)}, \Theta^{(2)}} -L(\Theta^{(1)}, \Theta^{(2)}) + \lambda_1 (||\Theta^{(1)}||_1 + ||\Theta^{(2)}||_1)$$

$$+ \lambda_2 \sum_{i=1}^{p} \left( ||\Theta^{(1)}_{i}||_2^2 + ||\Theta^{(2)}_{i}||_2^2 \right) ||V_i||_q$$

s.t. $\Theta^{(1)} - \Theta^{(2)} = V + V^T$

where $\sum_{i=1}^{p} \left( ||\Theta^{(1)}_{i}||_2^2 + ||\Theta^{(2)}_{i}||_2^2 \right) ||V_i||_q$ is the importance penalty and $w_j = ||\Theta^{(1)}_{j}||_2^2 + ||\Theta^{(2)}_{j}||_2^2$ is the importance weight of gene $j$. We followed PNJGL and decomposed the differential matrix symmetrically so that $V$ is not necessarily symmetric, and such decomposition will make the differential matrix easier to present the structure with hubs (Mohan et al., 2014). When the precision matrices $\hat{\Theta}^{(1)}$ and $\hat{\Theta}^{(2)}$ are adjacency matrices, i.e. binary matrices, the importance weight $w_j$ represents the sum of the degrees of gene $j$ in each of the two networks. The vital genes are often hubs in the networks and therefore have large importance penalties. According to our assumptions, these genes are penalized to involve the differential edges between two states. In brief, IPJGL adjusts the magnitude of its second penalty term according to the importance weights $\{w_j\}_{j=1..p}$ of genes which are adaptively updated from the estimated networks of two states. With the differentiated importance penalties, IPJGL gives more chances to reveal those less important genes involved in the differential network, respectively. For example, the degree of gene 4 is slightly increased while gene 2 totally vanishes in the result of IPJGL (Fig. 1). If there is strong evidence from the data that a gene is significantly changed between two states, the gene still can be revealed even with high importance penalty. If the change of interactions containing a gene is small, even if the importance penalty is greatly reduced, the result of the gene will not be affected too much. Therefore, the major difference between IPJGL and other general methods is that our method pays more attention to less important genes with the significant changes between the two states.

We used the ADMM (alternating direction method of multipliers) algorithm to solve the IPJGL model. For details of the algorithm, please refer to Section 1 of Supplementary Materials.

2.3 Performance metric

It is well known that the partial correlation coefficient matrix $P = (p_{ij})_{p \times p}$ can be obtained from the precision matrix $\Theta_{p \times p} = \left\{\frac{\Theta_{ij}}{\sqrt{\Theta_{ii} \Theta_{jj}}} : i \neq j\right\}$. Obviously $\Theta_{ii} = 0 \iff p_{ii} = 0$, so the corresponding network structures of the two matrices are the same. Moreover, the value of the partial correlation coefficient reflects the strength of the conditional correlation, therefore we consider using $\Delta_{\text{cox}} = p^{(1)}_{ij} - p^{(2)}_{ij}$ as the real differential matrix, $I(\Delta_{\text{cox}} < 0)$ as the label, $\Delta = p^{(1)}_{ij} - p^{(2)}_{ij}$ as the predicted differential matrix, and $|\Delta| = |\Delta_{ij}|$ as the score to calculate AUPR (area under the precision-recall curve). We did not use AUC (area under the receiver operating characteristic curve) as an evaluation indicator because for the sparse network inference problem, the proportion of positive edges is extremely low, so there is not much information in the comparison of AUC.

Furthermore, we proposed a novel metric, the absolute value of the second principal component (APC2) to conduct gene-pair survival analysis, which can test whether the interaction between a pair of genes is significantly changed for individual samples. In general survival analysis, we often consider a single gene and group samples according to its expression level. However, many diseases are unlike single-gene survival analysis on any of these genes alone did not yield significant results.

Specifically, APC2 is the absolute value of the second principal component based on principal component analysis. We denote $(X^T X)_{n \times 2}$ as the expression values of a pair of genes on n samples, which is centered on $(0, 0)$. $X = U \Sigma V^T$ is the singular value decomposition of $X$, where $(U)_{2 \times 2}$ is the eigenvector matrix of $X^T X$, $(\Sigma)_{2 \times 2}$ is the singular value matrix, $(V)_{n \times 2}$ is the eigenvector matrix of $V^T X$. The principal components of $X$ can be obtained by $Z = V \Sigma$, where $Z$ is a n x 2 matrix: the first column $Z_1$ is the first principal component (PC1) and $Z_2$ is PC2. Therefore, the APC2 of sample $i$ is $|Z_{i2}|$. The first principal component can be considered as the most correlated direction in all samples, and the APC2 of a sample captures its deviation from the most correlated direction (Supplementary Figs S22–S24). In the gene-pair survival analysis, given a pair of genes, we grouped the samples according to their APC2 values. We divided samples into two groups: half of the samples with smaller APC2 are in the group ‘high correlation’, and the other is in the group ‘low correlation’. Using gene-pair survival analysis based on APC2, we were able to identify some gene pairs that were significantly associated with patient survival time, whereas single-gene survival analysis on any of these genes alone did not yield significant results.
3 Simulation experiments

3.1 Data generation

Barabasi and Albert (1999) showed that many biological networks have a scale-free structure and proposed a BA algorithm for simulating the scale-free networks. In this article, we first used the BA algorithm to generate a scale-free network with the adjacency matrix \( A^{(1)} \), where \( p \) refers to the total number of genes. Then we generated a weight matrix \( U^{(1)} \) of the same size as \( A^{(1)} \), each of whose entry is sampled from the piecewise uniform distribution \( \left[-u_{\text{max}}, -u_{\text{min}} \right] \cup \left[u_{\text{min}}, u_{\text{max}} \right] \), and combine it with \( A^{(1)} \) to obtain \( \Theta^{(1)} \), i.e. \( \Theta^{(1)} = A^{(1)} \cup U^{(1)} \), where \( \circ \) refers to Hadamard product. This experiment takes \( u_{\text{max}} = 0.8 \), \( u_{\text{min}} = 0.3 \).

Second, we need to decide which genes are mutated. According to the assumptions, the importance of genes is correlated with their degrees in the network. Therefore, we divided the genes into three sets according to their degrees in the adjacency matrix \( A \) (because it is an undirected graph, we will generally much smaller than the probability of losing an existed interaction, i.e. \( \Theta_{ij} = 0 \)). We used the default parameter of each method (Fig. 2 and Supplementary Figs S2–S5). We used the selected \( \Theta^{(1)} \) and \( \Theta^{(2)} \) have at least 10% edges of the complete graph, and got \( \lambda_1 \) space \( \Lambda_1 = \{0, 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, \ldots, \lambda_2\text{max}\} \). Then, we looked for \( \lambda_{2\text{max}} \) (fixed \( \lambda_1 = 0 \)) to make both \( \Theta^{(1)} \) and \( \Theta^{(2)} \) have at least 2% edges (max edges of the true networks) of the complete graph, and got \( \lambda_2 \) space \( \Lambda_2 = \{0, 0.1\times \lambda_{2\text{max}}, 0.2\times \lambda_{2\text{max}}, \ldots, 0.9\times \lambda_{2\text{max}}, \lambda_{2\text{max}}\} \). Finally, we got the overall parameter space \( \Lambda = \Lambda_1 \times \Lambda_2 \), which can cover most network sparsity. For each dataset \( D_p \), \( r_e, r_d \) generated by the different combination of simulation parameters, we repeated 20 times and then recorded the \( (\lambda_1, \lambda_2) \) once if its AUPR is greater than 0.95 times the maximum AUPR of all parameter combinations. Finally, we obtained the frequency heatmap and select the best parameter combination \((\lambda_1^*, \lambda_2^*)\) with the highest frequency as the default parameter of each method (Fig. 2 and Supplementary Figs S2–S4).

Table 1. The distribution of mutated genes for five differential modes

|     | diffmode | 1   | 2   | 3   | 4   | 5   |
|-----|----------|-----|-----|-----|-----|-----|
| \(G_{\text{Hgb}}\) | m       | 0   | 0   | 0.5m| 0.1m|     |
| \(G_{\text{Middle}}\) | 0      | m   | 0   | 0   | 0.4m|     |
| \(G_{\text{Low}}\)  | 0      | 0   | m   | 0.5m|     |     |

Fig. 2. Frequency heatmap of parameter settings for IPJGL (\( q = 2 \)). X-axis refers to the \( \lambda_1^* \) (fixed \( \lambda_2 = 0 \)) of the parameter space \( \Lambda_1 \), Y-axis refers to the \( \lambda_1^* \) in the parameter space \( \Lambda_1 \). The max value of a cell is 1.

3.2 Parameter selection

All the methods to be compared use two parameters \( \lambda_1, \lambda_2 \). Therefore, we adopted a more strict framework for parameter selection following Tian et al. (2016) and Mohan et al. (2014). To compare the performance under each parameter combination in a comprehensive and detailed manner, we first determined the parameter space \( \Lambda \) for each method. We looked for \( \lambda_{1\text{max}} \) (fixed \( \lambda_2 = 0 \)) to make both \( \Theta^{(1)} \) and \( \Theta^{(2)} \) have at least 10% edges of the complete graph, and got \( \lambda_1 \) space \( \Lambda_1 = \{0, 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 30, 32, 34, 36, 38, \ldots, \lambda_{2\text{max}}\} \). Then, we looked for \( \lambda_{2\text{max}} \) (fixed \( \lambda_1 = 0 \)) to make \( \lambda_1 \) have at least 2% edges (max edges of the true networks) of the complete graph, and got \( \lambda_2 \) space \( \Lambda_2 = \{0, 0.1\times \lambda_{2\text{max}}, 0.2\times \lambda_{2\text{max}}, \ldots, 0.9\times \lambda_{2\text{max}}, \lambda_{2\text{max}}\} \). Finally, we got the overall parameter space \( \Lambda = \Lambda_1 \times \Lambda_2 \), which can cover most network sparsity. For each dataset \( D_p \), \( r_e, r_d \) generated by the different combination of simulation parameters, we repeated 20 times and then recorded the \( (\lambda_1, \lambda_2) \) once if its AUPR is greater than 0.95 times the maximum AUPR of all parameter combinations. Finally, we obtained the frequency heatmap and select the best parameter combination \((\lambda_1^*, \lambda_2^*)\) with the highest frequency as the default parameter of each method (Fig. 2 and Supplementary Figs S2–S4).

3.3 Simulation results

To show the effect of the proposed importance penalty, we compared IPJGL with several state-of-the-art methods JGL (Danaher et al., 2014), CNJGL, PNJGL (Mohan et al., 2014), because all these methods use similar GGMs to infer differential networks. We set \( n = 25, 50, 100 \), \( p = 100 \), \( q = 2 \), \( m = 10 \) to satisfy the small samples assumption. It can be seen from the equation that when \( q = 1 \), PNJGL is equivalent to JGL, and in our experiments, \( q = 2 \) always outperformed \( q = 1 \). Therefore, to show the results more clearly, we only show the results with \( q = 2 \) in the article. We used the selected default parameter \((\lambda_1^*, \lambda_2^*)\) for each method to predict differential networks. Each method kept the top 1000 to 50 edges in the differential network as the prediction results, and we repeated 50 times in each case and calculated AUPR (Supplementary Figs S5–S19). To evaluate the overall performance of each method on different numbers of kept edges, we show the average AUPR of each method. Here, we only show the case \( n = 100 \) (Fig. 3), the rest we have put in...
colorctal and breast cancer pathway including itself, which include 1034 and 958 genes, respectively. Next, we filtered out genes that did not express in at least 80% of the samples. Finally, we got 1009 and 953 genes, which are still relatively large and challenging gene sets for differential network inference problems, for colorectal and breast cancer, respectively. We used $\log(x + 1)$ to scale the data, and then standardize it as the final input.

4.2 Comparison with single-gene-based analyses

As a widely used technique, differential expression analysis (DEA) typically focuses on a single gene and then compares the difference between the two states for each gene, e.g. in expression levels. DEA is not only fast but also simple, straightforward and easy to understand. However, sometimes some genes are not significantly differentially expressed but their patterns of interacting with other genes are dramatically altered (de la Fuente, 2010). Fortunately, DiNA can determine which interactions between genes have produced changes, and if most interactions of some gene have changed, that gene can be considered the main factor in the changes between states. To investigate the power of DiNA, we are particularly interested in the results that were only discovered by DiNA method but not found by the traditional method of DEA.

We first applied the DiNA methods with the default parameters to the real dataset to obtain the initial differential networks, and then keep the top k edges as the result (denoted as ‘methodk’, e.g. the network generated by IPJGL with the top 500 edges is IPJGL500). Next, we used the ‘DESeq2’ package (Love et al., 2014) in R to perform DEA. The inputs to ‘DESeq2’ are the gene expression matrices in normal and disease states. The output of ‘DESeq2’, which we used, is a list of P-values for the genes. If the P-value for a gene is < 0.05, we consider the gene to be significantly differential in expression level between the two states. We then marked the insignificant genes (P-value > 0.05) with red color, which means those genes are less likely found by DEA but discovered by DiNA (Fig. 4).

To better exhibit the differential gene pairs discovered by DiNA, we used the metric APC2 as described in Section 2.3 to conduct gene-pair survival analysis on the gene pairs connected in differential networks. We downloaded the clinical data of TCGA and performed survival analysis using the R package ‘survminer’ on the single genes and the gene pairs. We fitted the data with the Kaplan–Meier model (Kaplan and Meier, 1958), which determines whether a difference in a factor leads to a difference in the probability of patient survival. For the single-gene survival analysis, the factor is the expression level between the two states. We then marked the insignificant genes (P-value > 0.05) with red color, which means those genes are less likely found by DEA but discovered by DiNA (Fig. 4).

4 Applications in TCGA data

4.1 Datasets preparation

Colorectal cancer and breast cancer are common and plague the normal life of many people. To explore if our method can extract some insights from real-world data, we applied IPJGL to the transcriptome profiles downloaded from the TCGA project. The datasets of HTSeq counts are used in experiments. For colorectal cancer, which belongs to adenomas and adenocarcinomas, there are a total of 39 normal samples and 398 cancer samples according to annotations. For breast cancer, which belongs to ductal and lobular neoplasms, there are a total of 112 normal samples and 1066 cancer samples according to annotations.

Due to the huge number of genes, we performed the following screening. First, we selected 10 and 9 pathways related to KEGG’s
Cytoscape’s plugin ReatomeFIVz (Kusuma et al. 2003; Semenov et al., 2005) further validated the correlation between SOST’s neighbors and colorectal cancer. A lot of evidence was found to prove the relationship between its neighbors and colorectal cancer, which further validates the correlation between SOST and colorectal cancer (Section 6.3 of Supplementary Materials and Figs S26 and S27). We also compared the relevance of the gene sets in the differential networks obtained by different algorithms in the Reactome database. We found that genes picked by IPJGL have more interactions in Reactome than that by PNJGL when keeping the same edges as the output (Supplementary Fig. S28), which implies the gene set found by IPJGL is more closely related in the known colorectal cancer database.

### 4.3 Colorectal cancer analysis

We compared IPJGL with PNJGL for clarity of results on the real datasets because PNJGL is second only to our method in the previous simulation experiments. We kept the top 10 genes ranked by degree in method10, method50, method100, method500 (Table 2 and Supplementary Tables S3–S5). For example, top 10 genes in method500 for colorectal cancer are shown in Table 2, where genes that cannot be found by DEA but can be discovered by DiNA are marked in bold. We integrated all bold genes in these four tables as the key genes identified by the corresponding method.

### Table 2. Comparison of the top 10 genes in 500-edges differential networks for colorectal cancer

| Rank | Gene      | Degree | P-value | Rank | Gene      | Degree | P-value |
|------|-----------|--------|---------|------|-----------|--------|---------|
| 1    | TCL1B     | 15     | 0.030   | 17   | INHBC     | 16     | 0.009   |
| 2    | HJV       | 14     | 0.735   | 18   | DKK1      | 16     | 0.000   |
| 3    | PK6       | 12     | 0.010   | 19   | WNT7A     | 16     | 0.000   |
| 4    | TUBA3C    | 12     | 0.071   | 20   | TUBA3C    | 14     | 0.071   |
| 5    | PPP3R2    | 11     | 0.635   | 21   | GHC1      | 14     | 0.000   |
| 6    | SOST      | 11     | 0.487   | 22   | INHBC     | 11     | 0.009   |
| 7    | TBL1Y     | 11     | 0.921   | 23   | FGF22     | 14     | 0.209   |
| 8    | TBL1Y     | 15     | 0.000   | 24   | FGF8      | 13     | 0.010   |

Note: The P-value is calculated using DESeq2. Genes with P-value >0.05 are marked in bold, which means they cannot be found by DEA but can be discovered by DiNA. If the degrees are the same, we order them by the sum of their interaction values.

We performed KEGG and GO enrichment analysis on genes in IPJGL500 of breast cancer. The X-axis (GeneRatio) represents the proportion of genes in the corresponding pathway that are covered by IPJGL500. The node size represents the number of genes in the intersection. The node color represents the adjusted P-value, where red indicates a smaller P-value. We successfully recover the breast cancer pathway using TCGA breast cancer data. The second significant pathway is the WNT signaling pathway, which is relative to a certain number of cancers.

### 4.4 Breast cancer analysis

We performed KEGG and GO enrichment analysis on genes in the differential networks with 50, 100 and 500 edges for breast cancer.
(Supplementary Figs S29 and S30). Because our candidate gene set is obtained from the KEGG pathways, we expected that the genes obtained by the algorithm should enrich the pathways related to breast cancer. The results confirmed that the genes obtained by IPJGL have strong functional enrichment properties, and only in the genes of IPJGL500, we obtained the expected results, as shown in Figure 5e. Although the dataset is relatively rough, the sample ratio is unbalanced, and the expression value may be biased, our method can still identify the corresponding pathway (breast cancer).

Further, we followed the same data analysis procedure in previous section to obtain key genes for breast cancer. We first retained all genes in the differential network method500 with P-value $>0.05$ in DESeq2. Then we performed the survival analysis for these genes to obtain the final set of key genes (Supplementary Fig. S31). Among them, there are 5 key genes obtained by IPJGL: SFRP5, FGF19, HE55, IL7R, RBBP8, and 3 key genes obtained by PNNJGL: SFRP5, FGFI9, HE55. The unique key gene RBBP8 identified by IPJGL is also called CTIF, whose low-level expression is associated with a poor prognosis in breast cancer (Wang et al., 2016), as shown in our survival analysis (Fig. 5f).

5 Conclusion

In this article, we developed a novel algorithm IPJGL for differential network inference by adding an importance regularization term in the GGM. The comprehensive simulation experiments of comparing the proposed method with several state-of-the-art differential network inference methods based on GGMs confirmed the advantages and validity of the new algorithm. We also clarified the difference between DiNA and DEA in that DEA focuses on individual genes, whereas DiNA focuses on gene interactions, which are more complex but can also reveal more information. For this purpose, we proposed a novel metric APC2 for evaluating the interaction between a pair of genes for individual samples, which can be used in the downstream analyses of DiNA such as the gene-pair survival analysis. Finally, by applying the new method on the TCGA datasets, we obtained some significant cancer genes, such as SOST and RBBP8 for colorectal cancer and breast cancer, respectively, and found a lot of evidence related to cancers. In a nutshell, we hope our algorithm, simulation studies, and gene-pair interaction measurement can help biological and medical researchers find more candidate key genes about various kinds of cancers, while there is still a lot of work that needs to do based on the assumption of gene importance.

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