Gene expression data analysis using Hellinger correlation in weighted gene co-expression networks (WGCNA)

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A R T I C L E   I N F O

Article history:
Received 13 March 2022
Received in revised form 9 July 2022
Accepted 9 July 2022
Available online 13 July 2022

Keywords:
WGCNA
Non-linear correlation
Alzheimer's disease
Hellinger correlation
GTEx
scRNA-seq

A B S T R A C T

Weighted gene co-expression network analysis (WGCNA) is used to detect clusters with highly correlated genes. Measurements of correlation most typically rely on linear relationships. However, a linear relationship does not always model pairwise functional-related dependence between genes. In this paper, we first compared 6 different correlation methods in their ability to capture complex dependence between genes in three different tissues. Next, we compared their gene-pairwise coefficient results and corresponding WGCNA results. Finally, we applied a recently proposed correlation method, Hellinger correlation, as a more sensitive correlation measurement in WGCNA. To test this method, we constructed gene networks containing co-expression gene modules from RNA-seq data of human frontal cortex from Alzheimer's disease patients. To test the generality, we also used a microarray data set from human frontal cortex, single cell RNA-seq data from human prefrontal cortex, RNA-seq data from human temporal cortex, and GTEx data from heart. The Hellinger correlation method captures essentially similar results as other linear correlations in WGCNA, but provides additional new functional relationships as exemplified by uncovering a link between inflammation and mitochondria function. We validated the network constructed with the microarray and single cell sequencing data sets and a RNA-seq dataset of temporal cortex. We observed that this new correlation method enables the detection of non-linear biologically meaningful relationships among genes robustly and provides a complementary new approach to WGCNA. Thus, the application of Hellinger correlation to WGCNA provides a more flexible correlation approach to modelling networks in gene expression analysis that uncovers novel network relationships.

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1. Introduction

Weighted gene co-expression network analysis (WGCNA) is a popular method for detecting and investigating highly correlated gene clusters based upon high-throughput sequencing expression datasets. These gene clusters may then act in coordination to facilitate specific biological processes [1–5]. The default correlation method of traditional WGCNA is Pearson's correlation, a standard measurement of linear correlation between two variables, measuring the extent of pairwise dependence between genes based on their expression level. However, linear correlation, or more generally monotone correlation, is not the only form of dependence (e.g. parabolic function, trigonometric function, etc.) [6]. Although dominated by several historical methods, such as Pearson's correlation, Spearman's correlation, and Biweight midcorrelation, new statistical methods quantifying dependence are constantly being explored. Mutual information (MI) quantifies complex dependence by measuring how much a random variable can be determined by knowing another [7]. Distance correlation is a new method that has the ability to detect both linear and non-linear dependence but has a higher priority for detecting linear dependence [8]. Hellinger correlation is a recently proposed estimator, which has better performance in characterizing general dependence. In contrast, Pearson's correlation and Biweight midcorrelation capture perfect linear dependence, and some rank-based measures such as Spearman's correlation only achieve their best performance in monotone dependence cases [9,10].

https://doi.org/10.1016/j.csbj.2022.07.018
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Alzheimer’s disease (AD) is the most common neurodegenerative disease that is pathologically characterized by β-amyloid (Aβ)-containing extracellular plaques and tau-containing intracellular neurofibrillary tangles, and clinically characterized by gradual cognitive decline and dementia [11–13]. Several hypotheses have been proposed to underlie AD including amyloidosis [14–17], tau pathology [18–20], mitochondria dysfunction [21–23], oxidative stress [23–25], and neuroinflammation [26–28], whereas the etiology remains elusive. Large-scale genome-wide association studies (GWAS) identified several risk genes of AD, such as mutations in APP (encoding amyloid precursor protein), AGRN (encoding agrin), LILRB2 (encoding leukocyte immunoglobulin-like receptor B2), GRN (encoding granulin), and APOE (apolipoprotein E) [29]. However, no single gene has been identified to explain the mechanism of most AD cases. Therefore, network-based analysis, representing a more comprehensive approach to recapitulate and investigate the pathogenic mechanism at the molecular level, is popular in understanding pathogenesis and discovering therapeutic targets in human disease [30–33].

In the present study, we demonstrate the insensitivity and insufficiency of calculating linear dependence (Pearson’s correlation, Spearman’s correlation, and Biweight midcorrelation) of gene expression to predict corresponding functional dependence. Subsequently, we applied 6 WGCNA methods: Pearson WGCNA and Spearman WGCNA, Biweight WGCNA, Distance WGCNA, MI WGCNA and Helinger WGCNA based on the corresponding correlation coefficients measuring gene dependence in RNA-seq human data sets including dorsolateral prefrontal cortex (DLPFC), temporal cortex (TC) and heart. The modules with highly correlated genes were compared. Finally, we focused on Helinger WGCNA results of DLPFC, and investigated the modules selected by the significant correlation coefficients measuring gene dependence in RNA-seq human data sets including dorsolateral prefrontal cortex (DLPFC), temporal cortex (TC) and heart. The modules with highly correlated genes were compared. Finally, we focused on Helinger WGCNA results of DLPFC, and investigated the modules selected by the significant correlation coefficients measuring gene dependence in RNA-seq human data sets including dorsolateral prefrontal cortex (DLPFC), temporal cortex (TC) and heart. The modules with highly correlated genes were compared. Finally, we focused on Helinger WGCNA results of DLPFC, and investigated the modules selected by the significant correlation coefficients measuring gene dependence in RNA-seq human data sets including dorsolateral prefrontal cortex (DLPFC), temporal cortex (TC) and heart.

2. Materials and methods

2.1. Brief introduction of different correlation coefficients

Let \( X_1 \) and \( X_2 \) be two continuous random variables.

2.1.1. Pearson’s correlation

In order to measure the dependence between \( X_1 \) and \( X_2 \), Pearson’s correlation quantifies the similarity between covariance \( \text{cov}(X_1, X_2) \) and the product of standard deviations \( \sigma_{X_1} \sigma_{X_2} \), defined by:

\[
\rho = \frac{\text{cov}(X_1, X_2)}{\sigma_{X_1} \sigma_{X_2}}
\]

2.1.2. Spearman’s rank correlation coefficient

The Spearman correlation is defined as the Pearson’s correlation coefficient between the rank of two variables:

\[
\rho = \frac{\text{cov}(\text{Rank}(X_1), \text{Rank}(X_2))}{\sigma_{\text{Rank}(X_1)} \sigma_{\text{Rank}(X_2)}}
\]

Compared with Pearson’s correlation, Spearman correlation is less sensitive to outliers. However, when variables are translated into rank, detailed information is lost. Therefore, Spearman’s rank is less powerful than Pearson’s correlation when the data is normally distributed.

2.1.3. Biweight midcorrelation

It is defined as:

\[
bicor(x_1, x_2) = \frac{\sum_{i=1}^{n}(x_{1i} - \text{med}(x_1))(x_{2i} - \text{med}(x_2))w_{ij}^{(a)}}{\sqrt{\sum_{i=1}^{n}(x_{1i} - \text{med}(x_1))^2} \sqrt{\sum_{i=1}^{n}(x_{2i} - \text{med}(x_2))^2}}
\]

where:

\[
w_{ij}^{(a)} = (1 - u_i^2)^2 \frac{I(1 - |u_i|)}{9 \text{mad}(x)}
\]

\( \text{med}(x) \) is the median of \( x \), and \( \text{mad}(x) \) is the median absolute deviation of \( x \).

As a median-based correlation method, biweight correlation is more robust to outliers than Pearson’s correlation, and without losing excessive information compared with Spearman correlation.

2.1.4. Distance correlation

The representation of Distance correlation is analogous to methods mentioned above. But instead of sample moments (e.g., variance), distance correlation considers certain Euclidean distances between sample elements which are defined as:

\[V(x) = \frac{1}{P} \sum_{i=1}^{P} A_{ik}^2 \]

\[A_{ik} = a_{ik} - \bar{a}_k - \bar{a}_i + \bar{a}
\]

\[a_{ik} = ||X_i - X_k||_p\]

And:

\[V(x, y) = \frac{1}{P} \sum_{i=1}^{P} A_{ik}^2 A_{il}^2\]

Then, the distance correlation is defined as:

\[R_d^2(X_1, X_2) = \frac{V^2(x, y)}{V^2(x) V^2(y)}\]

Rigorous proof shows that \( R_d^2(X_1, X_2) \) ranges from 0 to 1, and \( R_d^2(X_1, X_2) = 0 \) if and only if \( X_1 \) and \( X_2 \) are independent while \( R_d^2(X_1, X_2) = 1 \) if and only if \( X_1 = aX_2 + b \), which means perfect linear dependence (8). Therefore, distance correlation measures all types of possible relationships, including linear and non-linear dependence. But have higher priority for linear correlation.

2.1.5. Mutual information

\[\text{MI}(X_1, X_2) = \sum_{x_1, x_2} P_{X_1, X_2}(x_1, x_2) \log \frac{P_{X_1}(x_1)P_{X_2}(y)}{P_{X_1, X_2}(x_1, x_2)} = \int \int \frac{\phi(t)}{\bar{\phi}(t)} dF_1(x_1) dF_2(x_2)\]

\[\phi(t) = t \log t\]

It is easy to notice that \( \text{MI}(X_1, X_2) \) ranges from 0 to \( +\infty \). However, previous proof indicates that there are cases in which MI goes to infinite without a strong dependence between \( X_1 \) and \( X_2 \) [34]. In practical applications, a normalization value from 0 to 1 for MI is usually used for comparison purposes. In this paper, we define MI as:

\[\text{MI}_{\text{normalized}}(X_1, X_2) = \frac{\text{MI}(X_1, X_2)}{\max(\text{Entropy}(X_1), \text{Entropy}(X_2))}\]

Where: \( \text{Entropy}(X_1) = -\sum_{X_1} P(x_1) \log P(x_1) \)
MI reasonably and equally detects both linear and non-linear dependence. However, the power of MI is lower than other coefficients mentioned in this paper [35,36].

2.1.6. Hellinger correlation

In contrast, the Hellinger correlation ($\eta$) measures the dependence of $X_1, X_2$ by quantifying the similarity between their joint distribution $F_{12}$ and marginal production $F_1F_2$. Given by:

$$\eta = \left(\frac{2}{\mathcal{B}}\right)\left(\mathcal{B}^4 + (4 - 3\mathcal{B})^4 - 2\right)^{\frac{1}{4}}$$

where:

$$\mathcal{B} = 1 - \frac{1}{2}\int \left(\frac{dF_{12}(x_1, x_2)}{\sqrt{dF_1(x_1)dF_2(x_2)}} - 1\right)^2 dF_1(x_1)dF_2(x_2)$$

Detailed information and estimation procedures of $\eta$ are shown in [9]. Hellinger correlation ranges from 0 to 1 (from completely independent to perfect dependence) and has comparable power compared to the above-mentioned methods. Compared with MI information, Hellinger coefficient has higher statistical power and achieves highest value if and only if there are perfect dependent cases. At the same time, unlike distance correlation, it does not prioritize linear correlation.

In this paper, we used WGCNA package [2] to estimate Pearson correlation (complexity: O(n)), Spearman correlation (complexity: O(n)), Biweight correlation (complexity: O(n)), and MI (complexity: O(n^2)); energy package [37,38] to estimate distance correlation (complexity: O(n)); HellCor package [9] to estimate Hellinger correlation (complexity: O(n^2)). Based on the properties of different coefficients, we named coefficients only measuring linear correlation as linear coefficients (Pearson, Spearman, and Biweight) while named coefficients measuring both linear and non-linear correlation as dependence-based coefficients. Since dependence-based coefficients identify dependence instead of linear correlation with direction, we built undirected WGCNA graphs. Therefore, we took the absolute value of linear coefficients during gene co-expression network construction.

2.2. Data collection and preprocessing

Table 1 shows the description of RNA datasets curated for the analysis.

RNA-seq DLPFC dataset. RОСМАР RNA-seq normalized data (syn3505720) and clinical file (syn3191087) were downloaded from SYNAPSE [31]. We filtered out genes with FPKM less than 1 in more than 50% of samples. A variance-stabilizing transformation from package DESeq2 [42] was applied. We clustered the whole samples on their Euclidean distance to test their similarity, and one sample was chosen to represent each cluster as a newly constructed pseudo-cell. Finally, genes overlapped with the 5000 genes selected from RNA-seq DLPFC dataset were chosen for further analyses.

RNA-seq Heart dataset. Heart RNA-seq normalized data was downloaded from GTex database [41] and processed in the same manner as RNA-seq DLPFC dataset. The top 5000 genes with the highest standard deviation and 430 samples were selected for the rest of the analyses.

RNA-seq temporal cortex (TC) dataset. TC RNA-seq normalized data (syn8466815) and clinical file (syn8466814) were downloaded from SYNAPSE [40] and processed in the same manner as RNA-seq DLPFC dataset. Genes intersected with the 5000 genes selected from RNA-seq DLPFC dataset were chosen for further analyses.

Single-cell sequencing dataset. RОСМАР single-cell sequencing data (syn18485175) and metadata (syn3157322) were downloaded from SYNAPSE [39]. Single-cell raw data was processed by the package Seurat [44]. Specifically, we kept genes detected in no less than 3 cells and kept all cells with at least 200 detected genes. Outlier cells in quality metrics, including unique gene counts, the ratio of mitochondrial relative to endogenous RNAs, total gene counts, were filtered out as they might represent dead (or unqualified) or doublets (or multiplets) cells. For the rest of cells that passed the quality control, we carried out log-normalization with a scale factor 10000. Louvain algorithm in Seurat was implemented to detect clusters. The cell type of each cluster was identified by the corresponding marker genes provided by a previous publication [39]. Considering the sparsity of single-cell sequencing data, we applied scWGCNA [45], a package aggregating k nearest neighboring cells (neighboring cells within a cell-type-specific cluster) as a newly constructed pseudo-cell. Finally, genes overlapped with the 5000 genes selected from RNA-seq DLPFC dataset were obtained to set up cell-type-specific gene expression matrices of pseudo-cells, which were included to validate the preservation of modules from RNA-seq DLPFC dataset WGCNA results.

2.3. WGCNA and identification of significant modules

Different coefficients were calculated to measure the pairwise dependence between genes. To comply with scale-free topology criterion and the recommendations of WGCNA use, we chose appropriate soft-thresholding powers to convert the gene expression matrices to adjacency matrices. Then topology overlap matrices (TOM) were calculated by adjacency matrices [46]. We then use hierarchical clustering and dynamic tree cut method to identify gene clusters [2,47]. All steps of WGCNA with different coefficients are the same except for the calculation to measure gene dependence.

2.4. Network validation

**STRING database.** We built Protein-Protein networks based on the annotation from the STRING database (version = 11.5) [48]. To increase the credibility of validation, high confidence (score threshold = 900) was referred to in determining the connection between two proteins.

Normalized Mutual Information (NMI) [49] and adjusted rand index (ARI) [50]. NMI and ARI are two common statistics measuring the similarity between two data clustering results. NMI ranges from 0 to 1 and ARI ranges from $-1$ to 1, which corresponds to completely different to exactly the same.

Module preservation. We mainly used z-summary [51], a network preservation statistic aggregating multiple preservation statistics (3 density-based statistics and 3 connectivity-based statistics), to quantify the conservation of the co-expression network in another dataset.

2.5. Modules-phenotype association analysis and Bayesian network construction

Module eigengene was defined as the first principal component of a module gene expression matrix [2]. We identified the module of interest by the correlation strength between module eigengene and phenotypes, including pathologic stages (braak stage, CERAD
score) and clinical behaviors (Clinical diagnosis of cognitive status (dcfdx_jv), Final consensus cognitive diagnosis (cogdx)).

Markov chain Monte Carlo (MCMC) methods for structure learning and sampling of Bayesian networks were demonstrated to have better performance [52]. We used order MCMC algorithm in BiDAG, and 20 million iterations were applied [53].

2.6 Gene set enrichment analysis and cell-type-specific expression analysis

We implemented gene ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analysis by g: Profiler [54], and cell-type-specific expression analysis by CSEA tool [55].

2.7 Hub genes identification

We used cytoHubba [56] in Cytoscape [57] to identify hub genes in the protein–protein network. For networks with linked genes larger than 300, we identified the top 200 hub genes and showed related subnetworks, otherwise we displayed the entire PPI network.

3.3 Results

3.1 Comparison of different correlation coefficients

3.1.1 Coefficients measuring both linear and nonlinear association fit complex dependence

We hypothesized that there are complex relationships beyond linear correlation between genes in their expression in biological tissues. To test this hypothesis, we plotted several typical pairwise relationships of genes from the DLPFC RNA-seq dataset (Fig. 1A,C,E; Fig. 2A,C,E,G). We found that all correlation metrics are competent in accepting significant linear dependence and rejecting purely independent gene pairs, except that Pearson correlation is relatively more sensitive to outliers (Fig. 1B,D,F). They perform differently in detecting more complex dependence relationships (Fig. 2B,D,F,H). In these cases, dependence-based coefficients have a higher correlation value. We subsampled the data and show the receiver operating characteristic (ROC) curves [58] to evaluate the performance and robustness of different correlation coefficients. The results indicate that only dependence-based coefficients can consistently detect all kinds of complex dependence (Fig. 2B,D,F,H). To further verify the authenticity of these complex dependencv dependencies, we provide 6 coefficients of these gene pairs in 53 GTEx tissues (Supplementary Table 1) [41]. We found that the dependence of HBB and RCN2 (independent); CRABP2 and HS3ST2 (independent with outliers) are not significant in most of the tissues by most coefficients. The dependence of CYTB and ND1 (linear dependent); KPS28 and RPL36 (threshold); RAC2 and CYBB (complex dependent) are significant in most of the tissues by most coefficients. The dependence of GSTM1 and GSTM2 (dependent with outliers) is significant in most of the tissues by Hellinger correlation. However, in liver tissue, where GSTM1 and GSTM2 are highly expressed [59], the dependence is significant in all coefficients. The dependence of MYC and COX2 (power function) is significant in most brain tissues and other tissues, especially in the kidney. Interestingly, MYC and COX2 also show power function in those tissues.

3.1.2 Comparison of different coefficients

In Fig. 3, we show the edge-agreement results of the top 10 percent of gene pairs with the highest correlation values of each coefficient in the DLPFC dataset. As we expected from the statistical properties of each method, the results of dependence-based coefficients are more different from linear coefficients. Outlier insensitive methods have more overlap with each other (Spearman and Biweight). For dependence-based coefficients, MI and Hellinger are more likely to agree as they equally measure the linear and non-linear processes based on dependence. In contrast, distance correlation is more prone to give higher value to linear processes, so the results agree more with linear coefficients. Fig. 3B shows the number of gene pairs in Fig. 3A that can be found in the protein–protein interaction database (STRING database). It shows that many gene pairs with higher dependence-based coefficients have the potential to be functionally related. The corresponding figures of TC and heart are shown in Supplementary Fig. 1.2. These results highlight the necessity and advantage of applying a dependence-based coefficient to detect more comprehensive gene networks that linear coefficients might miss or ignore.

3.1.3 Comparison WGCNA results using different coefficients

To investigate the consistency and novelty of applying dependence-based coefficients in WGCNA, we compared their module results in cluster agreement, network preservation, and Module-wise comparison. The NMI and ARI suggest the cluster agreement level of WGCNA cluster results with different coefficients in DLPFC (Table 2), while the results of TC and heart are shown in Supplementary Tables 2 and 3, respectively. The results of dependence-based coefficients are more similar, which also happens in linear coefficient results. Network preservation results show that most modules are preserved in the network constructed by different coefficients (Supplementary Fig. 3A,5). However, there are still few modules identified by dependent-based WGCNA that are not consistently preserved in networks built by other methods. For example, the turquoise module detected by Hellinger WGCNA appears to be a collection of several distant gene clusters in other WGCNA methods, and its density and connectivity are less preserved (Supplementary Fig. 3A, Supplementary Fig. 6). For DLPFC RNA-seq dataset, we also quantified module preservation in an independent microarray DLPFC dataset from another study (Fig. 4). We found significant preservation evidence for 9 in Pearson, 10 in Spearman, 10 in Biweight, 7 in Distance, 8 in MI, and 6 in Hellinger, indicating the modules identified by WGCNA with different coefficients are consistent in another DLPFC study. These results show the feasibility of applying dependence-based coefficients in WGCNA as a complementary approach to the standard approach (e.g. Pearson’s correlation).

3.2 Investigation of Hellinger WGCNA modules

Here, we show the results of Hellinger WGCNA in DLPFC RNA-seq dataset, and investigate the novel network constructed by its dependence-based coefficients.

3.2.1 Modules related to AD pathologic and clinical phenotypes

We identified modules associated with pathologic and clinical features from two aspects: (1) correlation between module eigengenes and clinical phenotypes; (2) module-phenotype network by Bayesian network structure learning. We found the module blue, turquoise, green and red are highly related to braak stage (severity of neurofibrillary tangle), CERAD score (severity of neuritic plaques), and clinical cognitive diagnosis (Fig. 5A). Bayesian network indicated the relationship between these 4 selected modules and clinical phenotypes (Fig. 5B). GO functional and KEGG pathway enrichment analyses indicated that blue module related to oxidative phosphorylation and mitochondrial function (mainly genome-coding genes); turquoise module associated with mitochondrial function (mainly mitochondria-coding genes) and immunity; red and green modules are significant with synaptic vesicle cycle, neurotransmitter, and other neuron functions.

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Fig. 1. Gene-wise relationships observed in DLPFC data and corresponding ROC curves. The scatter plots with axes represent related genes’ expression levels (variance-stabilizing transformed FPKM). To construct the ROC curves, we calculated the gene pair coefficients of 50 samples sampling from DLPFC data sets. The sampling process was repeated 1000 times, and the proportion of rejecting the null hypotheses at different significance level are shown in the ROC. The null hypothesis indicates independence. The scatter plots and ROC curves of independent (A, B), independent with outliers (C, D), and linear dependent (E,F) gene-pairs are shown.
Table 1
Description of RNA datasets curated for the analysis. References for the datasets are shown in parentheses after the dataset. Abbreviations: DLPFC, dorsolateral prefrontal cortex; AD, Alzheimer’s disease.

| Dataset                           | Tissue                                      | Sample Size |
|-----------------------------------|---------------------------------------------|-------------|
| RNA-seq dataset (31)              | DLPFC Control: 235 AD: 406                 |             |
| Microarray dataset (30)           | DLPFC Control: 101 AD: 129                 |             |
| Single-cell sequencing dataset (39)| Prefrontal cortex (Brodmann area 10) Control: 24 AD: 24 |             |
| RNA-seq dataset (40)              | Temporal cortex AD: 80 Control: 71 Path age: 30 Progressive supranuclear palsy (PSP): 81 |             |
| RNA-seq dataset (41)              | Heart                                       | 430         |

Fig. 3. Venn plot of gene pairs with top 10 % highest correlation values of each coefficient in DLPFC RNA-seq dataset. (A) Specifically, the gene-pairs with coefficients: Biweight ≥ 0.62 Pearson ≥ 0.63 Spearman ≥ 0.62 Hellinger ≥ 0.74 MI ≥ 0.16 Distance ≥ 0.61 were chosen and shown in Venn plots. (B) The number of gene pairs in panel (A) that can be found in the protein–protein interaction database (STRING database).

Table 2
NMI and ARI between the clustering results of WGCNA constructed by different coefficients in DLPFC. NMI and ARI measure the similarity between two data clustering results. NMI ranges from 0 to 1 and ARI ranges from −1 to 1, which corresponds to completely different to exactly same.

|       | Pearson | Spearman | Biweight | Distance | MI | Hellinger |
|-------|---------|----------|----------|----------|----|-----------|
| ARI   |         |          |          |          |    |           |
| Pearson | 1       | 0.38     | 0.54     | 0.63     | 0.37 | 0.21      |
| Spearman | 0.38   | 1        | 0.55     | 0.44     | 0.29 | 0.15      |
| Biweight | 0.54   | 0.55     | 1        | 0.58     | 0.38 | 0.19      |
| Distance | 0.63   | 0.44     | 0.58     | 1        | 0.39 | 0.23      |
| MI     | 0.37    | 0.29     | 0.38     | 0.39     | 1   | 0.30      |
| Hellinger | 0.21   | 0.15     | 0.19     | 0.23     | 0.30 | 1         |

|       | Pearson | Spearman | Biweight | Distance | MI | Hellinger |
|-------|---------|----------|----------|----------|----|-----------|
| NMI   |         |          |          |          |    |           |
| Pearson | 1       | 0.56     | 0.61     | 0.67     | 0.43 | 0.40      |
| Spearman | 0.56   | 1        | 0.67     | 0.59     | 0.40 | 0.38      |
| Biweight | 0.61   | 0.67     | 1        | 0.62     | 0.44 | 0.39      |
| Distance | 0.67   | 0.59     | 0.62     | 1        | 0.45 | 0.42      |
| MI     | 0.43    | 0.40     | 0.44     | 0.45     | 1   | 0.47      |
| Hellinger | 0.40   | 0.38     | 0.39     | 0.42     | 0.47 | 1         |

Fig. 2. Gene-wise relationships observed in DLPFC data and corresponding ROC curves. The scatter plots with axes represent related genes’ expression levels (variance-stabilizing transformed FPKM). To construct the ROC curves, we calculated the gene pair coefficients of 50 samples sampling from DLPFC data sets. The sampling process was repeated 1000 times, and the proportion of rejecting the null hypotheses at different significance level was shown in the ROC. The null hypothesis indicates independence. The scatter plots and ROC curves of dependent with outliers (A, B), threshold (C, D), complex dependent (E, F), and power function (G, H) are shown.
3.2.2. Validation and investigation of module turquoise

As module turquoise is the main difference between Hellinger WGCNA and WGCNA with other coefficients, we further validated and investigated module turquoise’s structural and functional construction. We performed cell-type-specific expression analysis and found genes in the turquoise module are significantly enriched in oligodendrocytes, astrocytes, and immune cells. We investigated module preservation of module turquoise in an independent single-cell prefrontal cortex (Brodmann area 10) dataset, and Fig. 6 shows turquoise is highly preserved in oligodendrocytes and astrocytes. We ignored the module preservation test in immune cells because we did not detect enough immune cells in this single-cell dataset (Fig. 6).

The top 200 hub genes of module turquoise are summarized in Fig. 7.

Fig. 7 shows that Hellinger Correlation WGCNA constructs a network between mitochondria-coding genes and inflammation. It is worth noting that a similar network was also constructed in TC RNA-seq dataset (blue module), indicating this network might consistently exist in brain tissue (Supplementary Fig. 7). Interestingly, this module in TC is also not preserved in the network constructed by linear coefficients (Supplementary Fig. 4A). The interaction between inflammation and mitochondrial were proposed by many previous studies. Specifically, mitochondria plays an important role in immune pathways by releasing components, including mtRNA, mtROS, and related proteins, while many inflammatory processes affect mitochondria dynamics and functions. [60–63]. In addition, the dysfunction within the dependence between mitochondria and inflammation were hypothesized and demonstrated in neurodegenerative diseases [63–65]. Recently, new therapeutic approaches, targeting the crosstalk between mitochondria and inflammation, such as COX, PPAR-γ, NO synthases, were proposed for neurodegenerative diseases [66,67]. However, this mitochondria and inflammation relationship cannot be recapitulated by linear coefficients, as the dependence within some gene pairs beyond linear correlation (e.g. Fig. 2G,H in the DPFC RNA-seq dataset).

4. Discussion

This study illustrates the ability of Pearson’s correlation and other linear coefficients to predict pairwise biological dependence between genes based on gene expression level. We identified a thousand pairs of genes, of which the STRING database annotated biological dependencies. However, the statistical dependence of these pairwise genes is easily detected by Hellinger correlation rather than linear coefficients, indicating the importance of including non-linear correlation statistics in gene-wise dependence tests.

Whole-transcriptome analyses, such as RNA-seq, microarray, and single-cell sequencing are conventional methods to explore the mechanisms of complex diseases [68]. Given the cost of sequencing experiments and inaccessibility of biopsy tissues, especially with large sample sizes of human tissues, more comprehensive analysis is needed to find disease mechanisms and therapeutic targets. We proposed integrating Hellinger Correlation as an alternative option of Pearson’s correlation in WGCNA analysis. In Hellinger WGCNA, the turquoise module, including a connection between mitochondria-coding genes and inflammation, was uniquely constructed. We verified the preservation of Hellinger
turquoise in an independent microarray study. The lack of preservation was later confirmed due to some turquoise hub genes (mainly mitochondrial coding genes) missing in the microarray dataset. Therefore, we further performed cell-type-specific expression analysis and found Hellinger turquoise module was enriched and preserved in astrocytes and oligodendrocytes. In fact, existing

Fig. 5. Modules-phenotype association analysis. (A) Correlation between the module eigengene and phenotypes. Numbers indicate correlation coefficients and p-values. (B) Bayesian network including modules and phenotypes.
studies suggest the cellular metabolism of the interaction between mitochondria function and inflammation pathway [69], while the related hypothesis and experimental demonstration was extended to neurodegenerative diseases [63–65]. Furthermore, we found MYC gene as an important hub gene linking mitochondrial coding genes and inflammation-related genes. As a proto-oncogene in

Fig. 6. Hellinger turquoise module preservation assessed in the single-cell sequencing data. (A) Results of cell-type-specific expression analysis of Hellinger turquoise preservation genes. Colors from yellow to red represent the enrichment significance, while the size of hexagon represents the cell-specific level (the smaller the hexagon, the more stringently cell type specific it is). (B) Hellinger modules preservation assessed in the astrocytes. The green dashed line (Z-summary = 10) marks the “strongly preserved” threshold and the blue dashed line (Z-summary = 2) marks the “moderately preserved” threshold. (C) Hellinger modules preservation assessed in the oligodendrocytes. The green dashed line (Z-summary = 10) marks the “strongly preserved” threshold and the blue dashed line (Z-summary = 2) marks the “moderately preserved” threshold. Z-summary quantifies the conservation of the co-expression network in another dataset. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
cancers. MYC controls the cell cycle, apoptosis, and metabolism [70]. In peripheral nervous system, axotomy elevates MYC, which promotes axon repair [71,72]. In the central nervous system, however, axotomy decreases MYC, and reduces axon repair [71]. This might be because maintaining and strengthening synaptic structure rather than cell regeneration is more necessary for the central nervous system during the evolution of mammals. Recently, many studies have reported the potential role of MYC in causing or accelerating AD through abnormal cell cycle re-entry [73], apoptosis induction [74], abnormal DNA synthesis [75], and other metabolism processes [76–78]. In addition, strong MYC expression was detected in AD astrocytes, where we demonstrated the preservation of the Hellinger turquoise module [79]. Therefore, the detailed biological function of the Hellinger turquoise module might provide an exciting opportunity to advance our knowledge of the AD mechanism. In the Bayesian network, 3 other Hellinger modules: blue, red, and green, were related to the AD progression from incipience to terminal. Correlation significance with phenotypes and enrichment analysis results substantiated their relationship with AD. However, since the minor differences between these Hellinger modules and the corresponding modules constructed by other coefficients, we did not discuss them in detail, but the significance of these modules is still worth noting.

Although the default function of WGCNA package is Pearson's correlation, other methods of calculating correlations can also be applied flexibly. The R [66] source code of combining Hellinger correlation to WGCNA is provided in the Supplementary text. The application of non-linear dependence rather than linear dependence in weighted gene co-expression network analysis has many advantages at the biological level. Non-linear dependence is better at modeling dosing-related saturation or threshold interaction in protein-binding processes and signaling pathways [67]. In addition, the existence of outliers or missing values in high-throughput sequencing data, especially for low expressed genes, affects the linear relationship between gene pairs. Therefore, non-linear correlation network is more appropriate to detect the complex biological gene-wise interaction and technical variation of high-throughput sequencing data, which provides a comprehensive basis for detecting hub genes as therapeutic targets or biomarkers. In this work, we only applied this non-linear dependence related WGCNA in the AD dataset, and more attempts in other datasets or diseases whose mechanisms are more clearly defined could better demonstrate the importance of this approach. In addition, we found Hellinger correlation is more sensitive than Pearson correlation and the mean connectivity of Hellinger network higher. Therefore, Hellinger network is easier to detect with less clusters than Pearson network with the premise of scale free topology.

While we were preparing this manuscript, Hou et al [80] published an approach that applies distance correlation in WGCNA and demonstrated the ability of distance correlation in non-linear dependence detection, robust to outliers in microarray (macrophage and liver) and RNA-seq (cervical cancer and pancreatic cancer). The advantages of Hellinger correlation we recommend in this paper over distance correlation is that Hellinger correlation provides a fairer measure of linear and non-linear relationship while distance correlation takes its maximum value only in the case of the perfect linear relationship [8,9]. As Pearson correlation, distance correlation is only defined if variables have finite second moment, which is not required by Hellinger correlation [8,9]. However, both Hellinger correlation and distance correlation are dedicated to detecting dependencies and cannot get positive or negative correlation information compared to Pearson correlation. Optimization by using different correlation measurements...
depending upon specific datasets or problems would be worth pursuing in the future.

In summary, we applied Hellinger correlation in quantifying the dependence between gene pairs in WGCNA analysis to Alzheimer’s disease data sets. The verification tests and downstream analyses results provide new insight into applying non-linear correlation statistics to construct gene networks. Such an application should complement current methods to obtain a more comprehensive understanding of biological processes underlying complex diseases.

CRediT authorship contribution statement

Tianjiao Zhang: Conceptualization, Data curation, Methodology, Formal analysis, Writing – original draft. Garry Wong: Conceptualization, Funding acquisition, Resources, Supervision, Writing – review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

We thank members of the Wong lab for discussion and critical reading and suggestions for the manuscript. Haibin Zhu is gratefully acknowledged for help and advice in running MATLAB source code. This work was supported in part by grant MYRG2020-00213-FHS from the Faculty of Health Sciences, University of Macau.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.csbj.2022.07.018.

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