Prediction of seed gene function in progressive diabetic neuropathy by a network-based inference method

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Abstract. Guilt by association (GBA) algorithm has been widely used to statistically predict gene functions, and network-based approach increases the confidence and veracity of identifying molecular signatures for diseases. This work proposed a network-based GBA method by integrating the GBA algorithm and network, to identify seed gene functions for progressive diabetic neuropathy (PDN). The inference of predicting seed gene functions comprised of three steps: i) Preparing gene lists and sets; ii) constructing a co-expression matrix (CEM) on gene lists by Spearman correlation coefficient (SCC) method and iii) predicting gene functions by GBA algorithm. Ultimately, seed gene functions were selected according to the area under the receiver operating characteristics curve (AUC) index. A total of 79 differentially expressed genes (DEGs) and 40 background gene ontology (GO) terms were regarded as gene lists and sets for the subsequent analyses, respectively. The predicted results obtained from the network-based GBA approach showed that 27.5% of all gene sets had a good classified performance with AUC >0.5. Most significantly, 3 gene sets with AUC >0.6 were denoted as seed gene functions for PDN, including binding, molecular function and regulation of the metabolic process. In summary, we predicted 3 seed gene functions for PDN compared with non-progressors utilizing network-based GBA algorithm. The findings provide insights to reveal pathological and molecular mechanism underlying PDN.

Introduction

Diabetic neuropathy (DN) is the most common and troublesome complication of diabetes, characterized by the gradual loss of peripheral axons, resulting in diminished sensation, pain, and eventual complete loss of sensation (1). Pathologically, it is the culprit of a series of interrelated metabolic abnormalities with insulin deficiency and hyperglycemia (2). Importantly, DN affects up to 60-70% of diabetics, leading to the highest morbidity and mortality rates and to a huge economic burden of diabetes care (3,4). However, in addition to controlling blood glucose levels, no effective treatment options have been found to prevent, slow or reverse the progression of DN and are not always achievable even in alert patients (5). Besides, a good understanding of pathological and molecular mechanism underlying DN might give help to explore effective therapy of this complicated disease.

The difference of gene expression levels could reflect the propensity of many diseases, and thus identifying gene functions has been an effective way to reveal the pathological mechanism of a disease at molecular level (6). ELAVL3, as a member of the Elavl family, is known as a neuronal Elavl because it is expressed in peripheral and central neurons throughout development (7). Ogawa et al hold the view that ELAVL3 is essential for maintaining Purkinje neuron axons (8). It has been reported that ELAVL3 regulates neuronal polarity via alternative splicing of embryo-specific exon in AnkyrinG (9). There is some evidence to suggest that Molecule Interacting with CasL-Like1 (MICALL1), involved in pre-cytokinetic events, acts as a membrane hub on tubular recycling endosomes (10). According to recent reports, tubular recirculating endosome biogenesis was regulated through p53-MICALL1 pathway (11). HEY2, as a member of hairy-related basic helix-loop-helix transcription factor subfamily, is involved in boundary formation and cell fate determination (12). Recent research has indicated that miR-98 activated the Notch signaling pathway by binding to HEY2 in Alzheimer's disease mice to improve mitochondrial dysfunction and oxidative stress (13). Our research results agree with previous studies, suggesting that ELAVL3, MICALL1, HEY2 genes have effects on neurological diseases. It has been demonstrated that by using variants based on the guilty susceptibility (GBA) algorithm, gene function prediction can

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be performed with very high statistical confidence assuming that the associations in the genetic data is necessary to establish culpability (14). Although various of techniques have been proposed for purpose of extending the GBA to indirect connections, only slight effectiveness has been discovered (15-18).

In the present study, a new method was proposed to predict seed gene functions for progressive DN (PDN) patients, by integrating the GBA algorithm and network-based method. To achieve this goal, gene expression data and gene ontology (GO) annotations were collected from the public databases. Differentially expressed genes (DEGs) were identified as gene lists and background GO terms were extracted as gene sets. The co-expression matrix (CEM) was constructed on gene lists by Spearman correlation coefficient (SCC) method.

Materials and methods

Preparing gene expression data. Gene expression data [GSE24290 (19)] for human DN were recruited from the public-free Gene Expression Omnibus (GEO). In brief, based on the density of myelinated fibers, GSE24290 divided the DN patient samples into two groups, progressors and non-progressors. In short, the patients in the progressive group lost ≥500 fibres/mm², and the patients in the non-progressive group lost ≤100 fibres/mm² over 52 weeks (20). By mapping these preprocessed probes to gene structures, a total of 10,570 genes were identified for PDN for subsequent analysis.

Collecting gene sets. All human GO annotations were prepared from the Gene Ontology Consortium (21,22). For purpose of making these retained GO terms more correlated to progressors, we took the intersections between DEGs and GO terms. If the number of DEGs for a GO term was smaller than 20, it was removed. Only GO terms including equal or more than 20 DEGs were reserved.

Identifying DEGs. DEGs between progressors and non-progressors were detected (23). The ImFit function implemented in Limma was utilized to perform linear fitting, empirical Bayes statistics and false discovery rate (FDR) calibration of the P-values on the data (24,25). The thresholds for DEGs were set as P<0.05 and $llog_2$ fold-changel >2.

Constructing CEM. To further investigate the correlations or interactions among DEGs, a CEM for them was constructed based on the SCC algorithm (26). If the SCC for a pair of genes was positive, it would indicate a positive linear correlation between the two genes. Similarly, a negative SCC refers to a negative relationship of the gene pair. In addition, the absolute SCC value of an interaction was denoted as its weight value. Furthermore, the higher the weight value across two genes, the stronger the interaction was, especially for 1. Otherwise, the 0 meant that there was no interaction between two genes. As a result, a CEN was constructed according to weight.

Network-based GBA algorithm. The GBA algorithm was combined with network to predict significant gene functions for progressors. Specifically, for a DEG in the CEM, we chose its adjacent genes to enrich to a GO term. Based on these GO

functional annotations, a multi-functionality (MF) score was assigned to each gene i in the CEM (14),

$$S(i) = \sum_{k \lor i \in GO_k} \frac{1}{Num_{in_k} * Num_{out_k}}$$

of which Num_{in_k} represented the number of genes within GO group *k*, whose weighting had the effect of giving contribution to a GO group; and Num_{out_k} was the number of genes outside GO group *k* in the CEM, whose weighting provided a corresponding weighting to genes not within the GO group. Note that we computed the AUC for evaluating classification performances between progressors and non-progressors. Here, for assessing the predictive power of machine learners in the support vector machine (SVM) model, AUC is an assessment of the accuracy of clinical classification performance (27). Most importantly, an AUC of 0.5 represents classification at chance levels, while an AUC of 1.0 represents a perfect classification. Thus we defined GO terms of AUC >0.6 as seed gene functions for PDN patients in the present report.

Results

Gene lists. In this report, based on 10,570 genes in GSE24290, we identified 79 DEGs between progressors and non-progressors by Limma package when setting the thresholds as P<0.05 and $llog_2$ fold-changel >2. All DEGs were ranked in ascending order of their P-values (Table I). We found that the most significant DEGs were *ELAVL3* (P=1.87E-02), *MICALL1* (P=2.51E-02), *HEY2* (P=3.42E-02), *PCDHB1* (P=3.69E-02) and *OR2S2* (P=4.81E-02). Importantly, the 79 DEGs were regarded as gene lists for further exploitation.

Gene sets. In the Gene Ontology Consortium database, there are 19,003 gene sets involved in 18,402 genes. To make these sets with a stable performance and more correlated to progressors, we chose GO terms from 20 to 1,000 in size and then took intersections between the reserved terms with the gene lists. Only gene sets containing intersected DEGs >20 were left in the subsequent analysis. Hereinafter, we defined the amount of intersected DEGs as the count value of this term. As a result, a total of 40 GO terms were determined (Table II), termed with background GO terms. There were three terms with Count >60, cellular component (GO:0005575, Count = 71), biological process (GO:0008150, Count = 67), and molecular function (GO:0003674, Count = 66). Furthermore, we calculated the DEGs by expressing the spectral data. According to the GO enrichment analysis, the enrichment of 79 DEGs in background GO terms was obtained (28). DEG annotated to a GO term, and if the gene is present in the GO term, the value is 1 (red), otherwise 0 (yellow). Finally, the heatmap was obtained from the heatmap package in the R language to analyze the above enrichment situation (Fig. 1). The result of this heatmap was in accordance with Table II. Ultimately, the background GO terms were the gene sets used.

CEM. With an attempt to investigate biological correlations among DEGs, a CEM with 79 nodes and 3,081 interactions were constructed based on the SCC, of which each interaction possessed a weight value to reveal the interacted strength

Rank	DEG	P-value	Rank	DEG	P-value
1	ELAVL3	0.018730149	41	FER1L4	0.237218317
2	MICALL1	0.025064168	42	TAS2R13	0.254492341
3	HEY2	0.034183011	43	SYN1	0.254519864
4	PCDHB1	0.036938421	44	KCNMB4	0.257387331
5	OR2S2	0.048112652	45	SLC35E1	0.258472445
6	CNTD2	0.055977842	46	PRX	0.267184163
7	DNAJB12	0.059213903	47	GIPC2	0.276805505
8	GPR21	0.061091712	48	RBM12B	0.298659702
9	PFN1	0.068351436	49	HACD1	0.302339336
10	ZMIZ2	0.073944462	50	NAALAD2	0.309726354
11	HSPA6	0.080467794	51	KLHL21	0.310864132
12	TRIM36	0.082922925	52	TMX2	0.316649651
13	MTHFD1	0.08313726	53	HEBP2	0.319262815
14	RPA4	0.091277628	54	RFC2	0.319345713
15	NIPAL2	0.101024558	55	CCDC134	0.327623954
16	HDDC2	0.106181304	56	MEPE	0.343481838
17	LIME1	0.107101383	57	TRAPPC9	0.374991751
18	MAPKAPK5-AS1	0.108867691	58	TXNRD1	0.384530053
19	CCDC186	0.116573636	59	CTDNEP1	0.386873156
20	FCF1	0.117569875	60	SH2D4A	0.39096557
21	PSMD5	0.118641952	61	ZNF512B	0.390996512
22	YTHDF3	0.130853807	62	TAGLN2	0.400497742
23	CDHR5	0.135952991	63	C10orf76	0.401347031
24	C6orf120	0.1416115	64	PIK3IP1	0.409666004
25	VWA7	0.144178178	65	C5orf30	0.420802441
26	ASF1A	0.147208457	66	PKNOX1	0.449555958
27	NELL2	0.162233252	67	DNAJB2	0.465398033
28	SRM	0.167545586	68	NCR2	0.474076273
29	SH3KBP1	0.176459493	69	GALNT8	0.475841943
30	USE1	0.178993287	70	IKZF3	0.480703371
31	INSIG1	0.179397171	71	SENP5	0.492341515
32	SSH3	0.18328187	72	POMGNT2	0.510089318
33	FAM66D	0.185333874	73	CALHM2	0.515258516
34	PGAP3	0.190879424	74	ATP8B4	0.5305693
35	PER2	0.192655455	75	GALNT10	0.543825962
36	CCS	0.193106321	76	WIZ	0.550465591
37	E2F8	0.200074614	77	FER	0.594006462
38	HTR1F	0.221840324	78	PITPNA	0.597706177
39	ZNF142	0.232301069	79	МҮОЗА	0.599378182
40	PPP1R3B	0.236879137			3.277270102

Table I. Gene list for progressive diabetic neuropathy (PDN).

between two genes. The weight distribution for interactions in this CEM showed the characteristic of good adjacent matrix that weights on its diagonal nearly equaled to 1, which suggested that the CEM had a good network scale property. In particular, an edge between C6orf120 and PGAP3 had the highest weight of 0.994. Moreover, to further evaluate the activities of genes in interactions of high weights, topological degree centrality analysis was conducted on all nodes in the CEM. The results showed that C6orf120 connected with 56 adjacent genes and thus had the highest degree. Subsequently, an assortativity coefficient was calculated to assess degree assortative mixing pattern extent. Consequently, the assortativity coefficient for the CEM was 0.829, indicating that the network had perfect assortative mixing patterns.

A sub-network was extracted from the CEM by selecting those interactions with weight >0.8 and visualized it by Cytoscape software (Fig. 2). There were 48 nodes and 330 edges in the sub-network. Among these nodes, *PGAP3*, *C6orf120* and *RBM12B* had higher degree than the others, which suggested their key roles in the PCN patients and

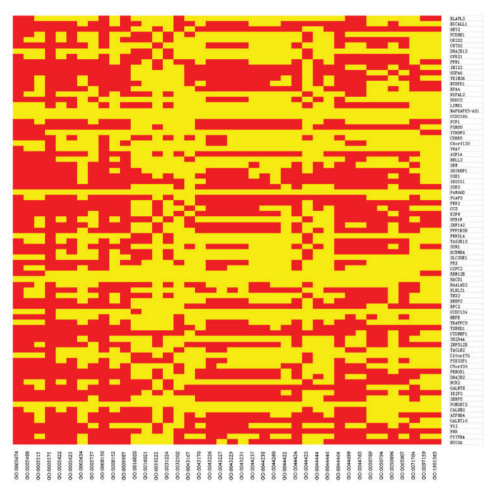


Figure 1. Heatmap for gene lists and sets. The ordinate, 79 differentially expressed genes (DEGs), the abscissa the 40 GO terms. Red, the association between the gene and the go term in sample channel. Yellow, no correlation between the gene and the go term in reference channel.

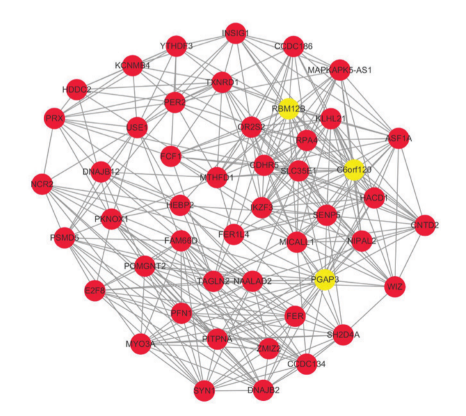


Figure 2. Sub-network of co-expression matrices (CEM). There were 48 nodes and 330 edges, of which the nodes were differentially expressed genes (DEGs) and the edges interactions between two DEGs with weight >0.8.

GO ID	GO term	Counts	GO ID	GO term	Counts
GO:0005575	Cellular component	71	GO:0008152	Metabolic process	36
GO:0008150	Biological process	67	GO:0071704	Organic substance metabolic process	34
GO:0003674	Molecular function	66	GO:0016020	Membrane	33
GO:0005623	Cell	58	GO:0044237	Cellular metabolic process	33
GO:0044464	Cell part	58	GO:0044238	Primary metabolic process	33
GO:0009987	Cellular process	57	GO:0043170	Macromolecule metabolic process	28
GO:0005488	Binding	55	GO:0044425	Membrane part	27
GO:0044699	Single-organism process	52	GO:0044444	Cytoplasmic part	27
GO:0043226	Organelle	49	GO:0044260	Cellular macromolecule metabolic process	26
GO:0044763	Single-organism cellular process	49	GO:0005634	Nucleus	25
GO:0005622	Intracellular	47	GO:0016021	Integral component of membrane	25
GO:0044424	Intracellular part	47	GO:0031224	Intrinsic component of membrane	25
GO:0043227	Membrane-bounded organelle	46	GO:0043167	Ion binding	24
GO:0065007	Biological regulation	43	GO:0050896	Response to stimulus	24
GO:0043229	Intracellular organelle	42	GO:0044422	Organelle part	23
GO:0005515	Protein binding	41	GO:0097159	Organic cyclic compound binding	23
GO:0043231	Intracellular membrane-bounded organelle	40	GO:1901363	Heterocyclic compound binding	23
GO:0050789	Regulation of biological process	39	GO:0044446	Intracellular organelle part	22
GO:0005737	Cytoplasm	38	GO:0019222	Regulation of metabolic process	21
GO:0050794	Regulation of cellular process	37	GO:0032502	Developmental process	21

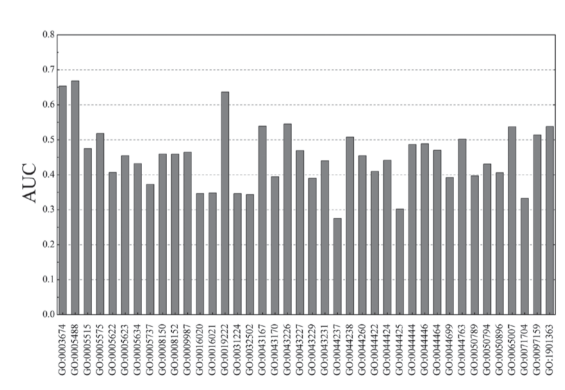


Table II. Gene sets for progressive diabetic neuropathy (PDN).



inferred that gene functions participated by them might act as critical processes in progressors.

Seed gene functions. The AUC index was applied to evaluate the classified performance on MF scores between progressors

and non-progressors by the 3-fold cross-validation. Generally, if an AUC for a gene or process was >0.5, it could be used to classify the case group from controls. In this report, the AUC for the 40 gene sets are shown in Fig. 3. There were 11 GO terms with AUC >0.5 in total. Thus, 11 of 40 (27.5%) gene sets

had a good classification performance between progressors and non-progressors of DN. Among them, 3 with AUC >0.6 were considered to be seed gene functions in progressors of DN. The seed gene functions were Binding (GO:0005488, AUC=0.668), Molecular function (GO:0003674, AUC=0.654), and Regulation of metabolic process (GO:0019222, AUC=0.636).

Discussion

In the present study, we predicted seed gene functions in PDN patients by using a network-based GBA method, since the network-based approach systematically investigate the molecular complexity of a particular disease (29) and identify potential signatures through bio-molecular networks rather than individual genes (30,31). An integration of co-expression network and the GBA algorithm provide a new manner to predict significant gene functions and reveal molecular mechanism underlying PDN.

On the basis of SCC method, a CEM for progressors was constructed on DEGs, and a sub-network of weight >0.8 was extracted from the CEM. Interestingly, we found that PGAP3, C6orf120 and RBM12B had high degree both in CEM and its sub-network, which indicated their importance in progressors. Taking RBM12B as an example, RBM12B (RNA binding motif protein 12B) is a protein coding gene that relates to functions of RNA binding, nucleic acid binding and nucleotide binding (32). It has been demonstrated that Rbm12b and *Rbm3* are mainly down-regulated and highly responsive to systemic hypoxia in mouse developing brain and placenta (33). This is the first time the key role of *RBM12B* in human PDN patients was uncovered. In addition, in our study, this gene was enriched in two seed gene functions, Binding and Molecular function. The possible inference was that the dys-regulation of *RBM12B* might disturb the normal functions of binding and lead to brain injury, even lesions of the nervous system.

Particularly, a total of 40 background GO terms were identified as gene sets for the current study. Subsequently, an MF score was assigned to each gene in the specific gene set, and then an AUC for each GO term was produced to assess the prediction performance between progressors and non-progressors. In consequence, 27.5% of all gene sets had a good classified performance with AUC >0.5. Most significantly, 3 gene sets with AUC >0.6 were denoted as seed gene functions for PDN, including Binding, Molecular function and Regulation of metabolic process. It is a common phenomenon that a given gene is present in one or more molecular functions (such as *RBM12B* described above), and two or more genes exhibit the same function.

Cameron *et al* reviewed that poor metabolic control was one of microvascular complications correlated to DN (34). Thus, it was important to improve the metabolic conditions that led to the pathology underlying peripheral DN, and the metabolic correction modified symptom control and clinical results (35). Moreover, metabolic dysfunction in experimental DN due to energy homeostasis and/or oxidative stress is limited to the sciatic nerve (36). Above all, we might conclude various metabolic processes play crucial roles in DP, and thus the regulation of metabolic process has become informative in PDN patients. In summary, we have predicted 3 seed gene functions for progressors of DP compared with non-progressors utilizing network-based GBA algorithm. The findings provide insights to reveal pathological and molecular mechanism underlying PDN. However, the expression data used in this work was recruited from the open access database, and the 3 seed gene functions still need to be validated.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

SSL and XBZ conceived the study and drafted the manuscript. XBZ, JMT and HRW acquired the data. SSL and THW analyzed the data and revised the manuscript. All the authors read and approved the final manuscript.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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