Integrated Bioinformatics Analysis of Hub Genes and Pathways Associated with a Compression Model of Spinal Cord Injury in Rats

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Background: Spinal cord injury (SCI) is a serious nervous system condition that can cause lifelong disability. The aim of this study was to identify potential molecular mechanisms and therapeutic targets for SCI.

Material/Methods: We constructed a weighted gene coexpression network and predicted which hub genes are involved in SCI. A compression model of SCI was established in 45 Sprague-Dawley rats, which were divided into 5 groups (n=9 per group): a sham operation group, and 1, 3, 5, and 7 days post-SCI groups. The spinal cord tissue on the injured site was harvested on 1, 3, 5, and 7 days after SCI and 3 days after surgery in the sham operation group. High-throughput sequencing was applied to investigate the expression profile of the mRNA in all samples. Differentially expressed genes were screened and included in weighted gene coexpression network analysis (WGCNA). Co-expressed modules and hub genes were identified by WGCNA. The biological functions of each module were investigated using the Gene Ontology and Kyoto Encyclopedia of Genes and Genomes databases.

Results: According to the RNA-seq data, a total of 1965 differentially expressed genes were screened, and WGCNA identified 10 coexpression modules and 5 hub genes. Module function analysis revealed that SCI was associated with immune response, cell division, neuron projection development, and collagen fibril organization.

Conclusions: Our study revealed dynamic changes in a variety of biological processes following SCI and identified 5 hub genes via WGCNA. These results provide insights into the molecular mechanisms and therapeutic targets of SCI.

MeSH Keywords: Genetic Association Studies • Pathologic Processes • Spinal Cord Injuries • Computational Biology

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Background

Traumatic spinal cord injury (SCI) is an important medical condition that causes disability in patients and imposes a heavy burden on patients’ families and the economy. A recent study estimated that there were 930 000 new cases of SCI worldwide, with an age-standardized incidence rate of 13 per 100 000 and 9.5 million years of life lived with disability. Ominously, SCI affects mostly young adults [1]. Compared with other central nervous system injuries, such as brain injury, SCI can result in greater long-term burden because patients with SCI usually survive the injury but have a poor life quality.

The pathophysiology of acute SCI involves primary (mechanical) and secondary mechanisms. Although both of these factors cause spinal cord damage and loss of motion, the secondary mechanisms warrant greater attention because they involve excessive inflammation, lipid peroxidation, inhibition of neurite growth, and delayed cell loss, which may be preventable or treatable [2,3]. Therefore, it is important to identify the key biological processes and hub genes involved in the secondary mechanisms of SCI.

High-throughput sequencing technology has become widely used in the field of SCI research. Transcriptome sequencing helped us identify many vital differentially expressed genes by comparing post-SCI and control samples.

Many molecular mechanisms were revealed by the functional analysis and in-depth bioinformatics analysis of these genes. However, the wealth of these abundant data, if only for differential expression analysis, was often underserved, reducing the value of high-throughput screening. Weighted gene co-expression network analysis (WGCNA) is a systematic biology approach that converts coexpression measures into link- age weight or overlapping topological measures and is widely used to explore the interactions between gene networks [4,5]. Generally, genes involved in the same pathway or functional complex show similar expression patterns and can be regarded as a module. WGCNA can be used to detect modules of highly correlated genes, summarize these clusters using the module eigengene or an intramodular hub gene, and correlate modules with other modules and external sample traits [6]. WGCNA is very suitable for monitoring the overall changes of modules with other modules and external sample traits [6].

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Most spinal cord injuries in the clinic feature a compressive component, while, at present, there has been no report of WGCNA being applied in the RNA sequencing result of spinal cord compressive injury. In this study, we used a compressive SCI model and sequenced the mRNAs of spinal cord samples at different times post-SCI. According to the WGCNA results, 10 co-expressed modules and 5 hub genes were identified, which may help to illustrate the molecular mechanisms of SCI and identify promising therapeutic targets.

Material and Methods

Animals and model of compressive SCI

A total of 45 male Sprague-Dawley rats (body weight 200 g to 250 g) were used in this study. The rats were obtained from Shanghai SLAC Laboratory Animal Co., Ltd. All rats were housed in a specific pathogen-free barrier system with a standard 12–12 h light/dark cycle at 23±1°C with free access to food and water in the laboratory animal center of Zhongshan Hospital, Shanghai, China. The rats were allowed to acclimate to the new system for 1 week before surgery was performed.

All animals were anesthetized by intraperitoneal injection of 5% chloral hydrate (0.6 mL/100 g body weight). During the operation, an electric blanket was used to maintain the body temperature of the rats, and ketoprofen (subcutaneous, 5 mg per kg body weight) was used for pain control. The rats were placed in independent cages after surgery. Each animal received penicillin by intramuscular injection (20 000 UI per 100 g body weight in 0.5 mL sterile saline) in the first 3 postoperative days to reduce infection. The animals’ bladders were expressed using the Credes maneuver (abdominopelvic compression) every 12 h to deal with SCI-related urinary retention until the rats were euthanized. This study was approved by the Committee of Ethics on Animal Experiments of Zhongshan Hospital.

The rats were randomly divided into 5 groups (n=9 per group): a sham operation group, and 1, 3, 5, and 7 days post-SCI groups. Within each group, 3 rats were used for RNA-seq and the others for histological examination (half for coronal section and half for transverse section). To prepare the clip compression model of SCI, the rats were anesthetized and underwent laminectomy at T8-9. Compressive injury was induced in the SCI groups by applying a 30 g aneurysm clip for 1 min, as previously described [7,8]. Rats in the sham operation group underwent laminectomy to expose the spinal cord, without clipping.

The animals in the SCI groups were euthanized at the corresponding time points, while the sham group was euthanized on the third postoperative day. The rats were euthanized by deep anesthetization via inhalation of isoflurane (4%). Tissue samples were subsequently quickly harvested, and the rats eventually died from the irreversible blood loss from tissue sampling.

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Histological examination

The spinal cords of the upper and lower 10 mm centered on the injured site were collected immediately and fixed in 4% paraformaldehyde, dehydrated in gradient alcohol, cleared in xylene, and embedded in paraffin. Then the embedded tissues were cut into 5-μm thick slices. The coronal section contained the full length of the longitudinal axis of the tissues, passing through the center of the spinal cord, and the transverse section was located at the site of the injury. These sections were then stained with hematoxylin and eosin (H&E) (Servicebio, Wuhan, China). The histological changes after SCI were recorded and analyzed under an optical imaging system (Leica Microsystems, Bensheim, Germany).

RNA isolation and sequencing

Spinal cord tissue from 5 mm around the epicenter of damage was harvested at the indicated times after inducing SCI and stored in liquid nitrogen. Total RNA was extracted using TRI Reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s protocol. The RNA concentration and quality were assessed using an automated electrophoresis system (Agilent 2100 Bioanalyzer, Agilent Technologies, Santa Clara, CA, USA). Subsequently, mRNA was purified from total RNA using oligo (dT)-attached magnetic beads for generating a cDNA library. The cDNA fragments were then sequenced on a sequencer (BGISEQ-500, Beijing Genomics Institute, Shenzhen, China). After quality control, raw data were mapped to the reference genome RAT6.0 and gene expression analysis was performed.

Identification of differentially expressed genes and WGCNA

Differentially expressed genes (DEGs) in the post-SCI groups were identified as genes that showed an increase in expression by >2 fold or a decrease by <0.5 fold relative to the sham group using the ‘limma’ package (v3.24.15) in R v3.4.1 [9]. The adjusted P values obtained by applying the Benjamini-Hochberg method for multiple testing correction (false discovery rate [FDR]) were considered significant when ≤0.05 [10]. Any post-SCI group with reads per kilobase per million (FPKM) >1 was used to construct the coexpressing modules. The gene coexpression network was constructed using the WGCNA package (v1.66) in R, as previously described [4,6]. All DEGs were included in the analysis. Briefly, the ‘pickSoftThreshold’ function was used to determine the correct power β after considering the smallest value to yield a free-scale topology. We selected 16 as the power value to obtain modules containing highly coexpressing genes. Using a dynamic tree-cutting algorithm and merging threshold function at 0.25, we identified 10 modules and summarized the expression profile of each module by its eigengenes. Expression trends of each module were plotted using the average fold change of the top 100 eigengenes, relative to the sham group, at each time point to reflect the overall change trend of each module.

Identification and Visualization of Hub Genes

We combined modules with similar expression trends to create new sets of genes. The top 150 connections among hub genes in each set were visualized into a weighted coexpression network using Cytoscape (v3.4.0) (http://www.cytoscape.org) [11]. Intramodular hub genes are usually located in the center of the module. The thickness of the edge was based on ‘weight’, which referred to the connection strength between 2 nodes (genes) in terms of the correlation value obtained from the ‘TOMsimilarity’ function in WGCNA. The Topological overlap matrix (TOM), which was obtained from ‘TOMsimilarity’, gave these values for the respective genes. This TOM value is always between 0 and 1. A higher value refers to a strong connection or coexpression of genes.

Gene enrichment analysis

The Gene Ontology (GO) [12,13] and Kyoto Encyclopedia of Genes and Genomes (KEGG, https://www.kegg.jp/) databases were used for functional annotation of specific modules using the Database for Annotation, Visualization and Integrated Discovery (DAVID) [14,15]. FDR <0.05 was considered as the threshold to indicate a statistically significant difference in the analysis.

Results

Morphological changes of SCI after compression

The animal model of SCI was established as shown in Figure 1A. All 45 rats received successful operations, with hindlimb paralysis and urinary retention observed in SCI rats and no obvious abnormalities observed in the sham group. Spinal cord disruption was visible after spinal cord compression (Figure 1B–1G). Images of the spinal cord and H&E-stained tissue sections revealed the extent of the pathological changes to the spinal cord (Figure 1H–1Q). Blood cells were visible 1 day after the injury, and tissue sections contained numerous red blood cells along the injury site with abundant inflammatory cells. Loss of the surrounding nerve tissue due to secondary injury was visible on day 3, at which time the blood had been absorbed. The tissue had undergone severe collapse, and scarring was evident. By day 5, the scar had grown in size with ongoing tissue repair, and the inflammation had diminished alongside the formation of a spinal cord cavity. By day 7, the transverse diameter of the spinal cord had decreased without obvious signs of clamp gross injury, but the injury site was mostly filled with scar tissue and a cavity.

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A total of 15 samples of spinal cord tissues from the injured rats (including 3 samples from the sham group) were homogenized and sequenced by BGISEQ-500 (Beijing Genomics Institute, China). Each group comprised 3 replicates. We searched for genes that were significantly upregulated or downregulated at any time compared with the sham group, as shown in Table 1, and identified a total of 1965 DEGs (Figure 2A–2D) for inclusion in the WGCNA.

WGCNA

We performed WGCNA to develop gene modules related to the pathology of SCI. For this screening, we set $\beta=16$ and identified the genes using the dynamicTreeCut algorithm in R. A total of 10 modules were split and each module was assigned a different color (Table 2). Heat maps were plotted to characterize the expression of these 10 modules. In this analysis, genes in the magenta and brown modules were downregulated on day 1 after injury, and tended to recover after this time. Genes in the blue, pink, green, grey, and red modules were downregulated on day 1 after injury, and then started to recover on day 3 and kept increasing after that. Generally, black and turquoise modules decreased, while the black module had a sudden decline on day 1 and then recovered. The expression of genes in the yellow module was upregulated on day 3, plateauing thereafter. The expression trends of genes in each module are shown in Figure 3A–3T.

Figure 1. Establishment of spinal cord injury after compression. (A) Diagram of compression model of spinal cord injury (SCI). (B–G) The figure demonstrated the visible signs of bleeding and spinal cord disruption after SCI. (H–Q) H&E staining showed the pathological changes at different time points (Sham, 1d, 3d, 5d, and 7d) after SCI in, coronal, and transverse sections (white arrows mark the injury site).
Biological functions analysis

We used DAVID to determine the biological functions of the genes in the 10 modules in terms of GO biological process terms and KEGG pathways. Table 2 provides an overview of the 10 modules, which contained the top 3 biological processes (BP) identified by GO analysis. Four of these modules were focused on the results of their KEGG pathway analysis, as illustrated in Figure 3U–3X. Genes in the largest module, turquoise, were related to chemical synaptic transmission, sensory perception of pain, and neurotransmitter secretion. These genes kept decreasing with time, especially at 5 days post-SCI, indicating they may have been regulated by injury-related factors, hindering neuronal remodeling. Genes in the blue module were related to immune function, being enriched with genes involved in inflammatory responses, innate immune responses, and adaptive immune responses. Pathway analysis also indicated that they were mainly related to inflammatory response. The blue module showed transient downregulation after injury followed by a gradual increase in expression, indicating that these genes may be involved in secondary damage, and that the immune system might play important roles in secondary injury and repair of the spinal cord following compressive injury. The GO-BP and KEGG analyses of the brown module indicated it was related to cell division, chromosome segregation, and mitotic nuclear division. These genes were markedly downregulated following injury but tended to recover after the acute stage. The green module showed a relationship with the extracellular matrix, collagen fibrils, and blood vessels according to GO-BP and KEGG analyses. The genes in the green module were suppressed on the third day, and then gradually increased and exceeded the baseline, which suggested a process of tissue reconstruction or glial scar formation.

Identification of hub genes

The modules with similar expression trends were firstly merged to obtain a larger gene network to help avoid the loss of marginal genes in some small modules, such as pink, black, and magenta. According to the expression results shown in Figure 3A, we merged the modules into 4 sets: set A (blue+green+grey+pink+red), set B (brown+magenta), set C (turquoise+black), and set D (yellow). Network analysis of the 4 sets allowed us to identify the driver genes via centralized hub analysis. We found that set A comprised 3 subsets, in which cytochrome P-450 (CYP) 2D1 (Cyp2d1) was the hub gene, while C2 and Cyth4 were the key nodes linking the subsets. For the other 3 sets, we found that the hub genes were Bub1b and Tpx2 for set B, sodium-potassium ATPase (Atp1a3) for set C, and Rpp9 and Pgams5 for set D (Figure 4). These hub genes were selected based on their degree of connection with other genes in the WGCNA network. The other major hub genes are listed in Table 3.

Discussion

RNA sequencing has been used extensively to identify potential therapeutic targets and prognostic markers in SCI. The pathological mechanism of SCI is complex and involves more than 20 potential pathological processes. To date, most studies that have performed RNA-seq in SCI have not fully exploited the data collected. Differential analysis and function enrichment...
Figure 2. Differential gene screening and weighted gene coexpression network analysis. (A) A total of 1965 differential genes were screened at different time points (Sham, 1d, 3d, 5d, and 7d) after spinal cord injury (SCI). (B, C) Analysis of network topology showed that it met the scale-free topology threshold of 0.8 when $\beta=16$. (D) The heatmap of topological matrix showed the correlation network, in which each color represented a gene module.
analysis were commonly used for mining RNA-seq data, but not enough for the massive data of high-throughput sequencing. In this study, we used WGCNA for a deeper analysis of RNA-seq; it integrated 1965 DEGs into 10 modules. Four relatively explainable modules were identified through the function enrichment analysis. They were the turquoise module related to pain or sense perception, the brown module related to cell division, the blue module related to immune response, and the green module related to extracellular matrix organization. By integrating similar modules into 4 gene sets, we identified Cyp2d1, Bub1b, Tpx2, Atp1a3, Rrp9, and Pgam5 as hub genes.

Inflammation is a double-edged sword for neurons in SCI. It is essential for the local tissue repairing but toxic if remained unresolved, which results in regeneration disorder [16]. In the present study, the increased expression of Pgam5 following SCI was observed. It was reported that Pgam5 could induce inflammasome activation and accelerate necroptosis and inflammation, which are involved in the early stage of secondary injury of SCI [17–19]. Therefore, it seems that Pgam5 may play a pathological role in SCI by accelerating necroptosis and inflammation. Meanwhile, the pathways of inflammatory response, innate immune response, and phagocytosis in the blue module were identified in our study, and we found they were suppressed within the first 3 days. These pathways were

| Module  | N   | Term                                      | Count | %     | FDR     |
|---------|-----|-------------------------------------------|-------|-------|---------|
| Turquoise | 679 | Chemical synaptic transmission          | 41    | 6.0   | 8.0E-23 |
|         |     | Sensory perception of pain                | 25    | 3.7   | 5.9E-17 |
|         |     | Neurotransmitter secretion                | 18    | 2.7   | 1.6E-16 |
| Blue    | 444 | Inflammatory response                     | 26    | 5.9   | 1.2E-08 |
|         |     | Innate immune response                    | 22    | 5.0   | 3.8E-07 |
|         |     | Phagocytosis, engulfment                  | 8     | 1.8   | 9.7E-06 |
| Brown   | 151 | Cell division                             | 24    | 15.9  | 1.0E-21 |
|         |     | Chromosome segregation                    | 17    | 11.3  | 4.7E-20 |
|         |     | Mitotic nuclear division                  | 16    | 10.6  | 4.9E-14 |
| Yellow  | 79  | rRNA processing                           | 5     | 6.3   | 1.5E-04 |
|         |     | tRNA methylation                          | 3     | 3.8   | 4.4E-03 |
|         |     | Ribosomal large subunit assembly          | 3     | 3.8   | 6.6E-03 |
| Green   | 63  | Collagen fibril organization              | 6     | 9.5   | 1.7E-07 |
|         |     | Blood vessel development                  | 6     | 9.5   | 2.9E-06 |
|         |     | Extracellular matrix organization         | 6     | 9.5   | 1.4E-05 |
| Red     | 30  | Innate immune response                    | 4     | 13.3  | 9.0E-03 |
|         |     | Cellular calcium ion homeostasis          | 3     | 10.0  | 1.1E-02 |
|         |     | High-density lipoprotein particle remodeling | 2   | 6.7   | 2.0E-02 |
| Pink    | 22  | Receptor-mediated endocytosis             | 2     | 9.1   | 1.0E-01 |
|         |     | Aging                                     | 2     | 9.1   | 2.9E-01 |
|         |     | Apoptotic process                         | 2     | 9.1   | 3.3E-01 |
| Black   | 22  | Neuron projection development             | 3     | 13.6  | 2.5E-03 |
|         |     | Protein localization to synapse           | 2     | 9.1   | 1.2E-02 |
|         |     | Signal transduction                       | 4     | 18.2  | 1.8E-02 |
| Magenta | 19  | Regulation of norepinephrine secretion    | 2     | 10.5  | 8.2E-03 |
|         |     | Adenosine receptor signaling pathway      | 2     | 10.5  | 1.0E-02 |
|         |     | Toll-like receptor signaling pathway      | 2     | 10.5  | 1.9E-02 |

Table 2. Overview of 10 modules with their top 3 GO-BP terms.

N – number of genes involved in the given module; Count – number of genes involved in the given term; % – percentage of involved genes/total genes; FDR – adjusted P values obtained applying the Benjamini-Hochberg method.
usually associated with microglia or macrophages, which may take part in the positive effects of inflammation and contribute to tissue repair after SCI [20].

The reconstruction of neural pathways is essential during the functional recovery after SCI. Cyp2d1, the hub gene of set A, has been reported to participate in a variety of neural activities [21]. According to our data of RNA-seq, Cyp2d1 was suppressed in the first 3 days and then gradually increased. A previous study reported a similar expression trend after traumatic brain injury [22]. Cyp2d1 was reported to be involved in the process of synaptic remodeling. Therefore, we speculated that Cyp2d1 may be a positive factor for nerve recovery. Another hub gene, Atp1a3, was persistently downregulated after SCI. This trend was also observed in a bioinformatics study that combined transcriptome and proteomics of contusive SCI in rats [23]. Several studies have demonstrated that mutations in Atp1a3 may induce persistent neurological dysfunction [24].

Figure 3. Expression trends and pathway enrichment of modules identified via WGCNA. (A–T) A total of 10 modules were split and each module was assigned a symbolic color. It was found that the modules of magenta and brown were sharply downregulated on the first day after injury, then tended to recover; the modules of blue, pink, green, grey, and red were downregulated after injury, and then rebounded upward 3 days after SCI; the modules of black and turquoise showed a continuous decrease after injury; the yellow module was sharply upregulated within 3 days after injury, and then tended to be flat. (U–X) The figures demonstrated the top 10 of pathway enrichment in blue, brown, green, and turquoise.
Hub genes were identified in 4 sets through weighted gene coexpression network analysis (WGCNA). It was found that set A had 3 subparts, Cyp2d1 was the hub gene of set A, which was the core of the network, and C2 and Cyth4 were the key nodes that linked the other 2 parts. The other 3 sets exhibited their respective aggregations. Bub1b and Tpx2 were hub genes of set B. Similarly, Atp1a3 was the hub gene of set C, while Rrp9 and Pgam5 were hub genes of set D. Line thickness represented connection weight. The size and color of the nodes represented the degree of connectivity.

Additionally, it was reported that Atp1a3 was downregulated in the freeze lesion microgyrus model, which suggests that Atp1a3 may be involved in neonatal cortical injury [25]. Atp1a3 is a sodium-potassium pump ATPase involved in the generation of electrical impulses by transporting neurotransmitters and calcium ions across cell membranes [23,26]. Disruption of the electrophysiological function of the spinal cord may be a major reason for the poor prognosis of SCI. Future studies of Atp1a3 may reveal more details about its role in SCI. Additionally, the results of the present study revealed that the expression of genes kept decreasing during the whole acute stage of SCI in the turquoise module, which was associated with neurotransmitter, synaptic transmission, and axon genesis. Further studies are required to explore the reasons behind these changes.
| Set   | Hub gene | Degree | Neighborhood connectivity | Topological coefficient | Betweenness centrality |
|-------|----------|--------|----------------------------|-------------------------|------------------------|
| **Set A** |          |        |                            |                         |                        |
|       | Cyp2d1   | 8      | 3.63                       | 0.25                    | 0.18                   |
|       | Gna15    | 6      | 4.83                       | 0.29                    | 0.46                   |
|       | Trem2    | 6      | 3.33                       | 0.25                    | 0.31                   |
|       | Cndp1    | 6      | 4.00                       | 0.38                    | 0.29                   |
|       | Fgd2     | 6      | 4.33                       | 0.32                    | 0.20                   |
| **Set B** |          |        |                            |                         |                        |
|       | Bub1B    | 26     | 14.92                      | 0.29                    | 0.10                   |
|       | Tpx2     | 26     | 14.73                      | 0.28                    | 0.12                   |
|       | Nusap1   | 22     | 13.55                      | 0.29                    | 0.20                   |
|       | Cdk1     | 22     | 16.32                      | 0.33                    | 0.04                   |
|       | Kifc1    | 21     | 15.67                      | 0.31                    | 0.06                   |
| **Set C** |          |        |                            |                         |                        |
|       | Atpl1a3  | 18     | 9.61                       | 0.21                    | 0.11                   |
|       | Rasgrf1  | 16     | 9.81                       | 0.20                    | 0.10                   |
|       | Sez6l2   | 14     | 9.64                       | 0.18                    | 0.21                   |
|       | Chga     | 14     | 9.86                       | 0.22                    | 0.09                   |
|       | Slc32a1  | 13     | 7.31                       | 0.17                    | 0.20                   |
| **Set D** |          |        |                            |                         |                        |
|       | Rtp9     | 19     | 8.58                       | 0.16                    | 0.19                   |
|       | Pgam5    | 18     | 8.39                       | 0.17                    | 0.13                   |
|       | Utp4     | 13     | 9.92                       | 0.20                    | 0.12                   |
|       | Spry2    | 12     | 7.42                       | 0.25                    | 0.06                   |
|       | Rnh1     | 11     | 10.82                      | 0.26                    | 0.06                   |

**Table 3.** Hub genes of module sets.

Glial scarring is a healing response but is unfavorable to axon growth in the long term [27]. Extracellular matrix organization is considered an important feature of scar formation [28]. In the present study, the green module showed a relationship with extracellular matrix, collagen fibril, and blood vessel development that suggested a process of tissue reconstruction, or glial scar formation. It was suppressed initially after SCI, and then gradually increased even beyond the baseline. This suggested that extracellular matrix failed to be activated in the early stage and failed to limit the expansion of inflammation, but was improperly activated in the later stage, causing obstruction of neural regeneration. This finding suggested that the precise regulation of extracellular matrix organization may be a potential therapeutic target for spinal cord injury.

Cell proliferation is a manifestation of tissue remodeling. The main pathways included in the brown module, such as cell division, chromosome segregation, and mitotic nuclear division, embodied this feature. We also identified Bub1B and TPX2 as hub genes of set B. Bub1B is considered essential for ensuring proper chromosome segregation, while TPX2 is suggested to redistribute from the nucleus to the apical process during the S-G2 transition [29,30]. These 2 genes belong to a family of mitosis-related genes [31,32]. Growing evidence indicates that mitosis plays an important role in tissue repair, but was long neglected [33,34]. The recovery of mitosis-related genes in the present study reflected the crucial role of mitosis in the recovery of other functional activities, which should be confirmed in future studies. However, it is suggested that cell cycle activation may contribute to secondary damage cascade after SCI [35]. Cell cycle proteins appeared to lead to apoptotic cell death and gliosis or microglial activation in SCI [36]. These contradictory findings reflected the complexity of SCI and needed further exploration.

WGCNA is an effective method for analyzing biological systems. In the present study, WGCNA found many hub genes and pathways that would not be noticed by routine differential analysis.
A small number of studies on SCI applied WGCNA. For example, Wang et al. used WGCNA in 2 open access array data and identified 3 hub genes including Rac2, Itgb2 and Tyrobp, which may play crucial roles in SCI via natural killer cell-mediated cytotoxicity [37]. Another study using WGCNA indicated that the effects of NT3-chitosan on neurite regeneration were mediated by the enhancement in neurogenesis and angiogenesis and reduced inflammatory responses [38]. Notwithstanding, the present study is not the first time that WGCNA has been applied on SCI; however, our work combined WGCNA with RNA-seq, which offered valuable insights into understanding the pathophysiological process of spinal cord injury.

This study had the following limitations. First, the spinal cord is a complex tissue with many cell types, and sequencing reflects expression changes in the whole tissue, rather than in specific cells. This may mistakenly attribute some functions to certain cells. More in-depth research in related types of cells is needed. The emerging topographic single-cell sequencing technology may solve this problem but it is too expensive and not mature in spinal cord tissue. Second, we did not validate the changes in gene expression in terms of protein expression. Due to the post-transcriptional regulation, the trend of protein expression may be different from that of mRNA.

Third, we obtained tissue samples up to 7 days after SCI, so future studies should include long-term observation following SCI. Finally, we failed to establish a hypothesis on the function of the last hub gene, Rp9, due to a lack of literature support. Future research is needed to evaluate its role in SCI.

Conclusions

In this study, we constructed a weighted gene coexpression network and predicted hub genes and essential biological processes in compressive spinal cord injury (SCI). Four essential modules were revealed to be associated with chemical synaptic transmission, immune response, cell division, and extracellular matrix organization. Moreover, Cyp2d1, Tubb1, Tpx2, Atp1a3, and Pgam5 were identified as hub genes. These findings may provide insights into the molecular mechanisms and therapeutic targets of SCI.

Conflicts of interest

None.

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