Identification of four differentially expressed genes associated with acute and chronic spinal cord injury based on bioinformatics data

Abstract

Complex pathological changes occur during the development of spinal cord injury (SCI), and determining the underlying molecular events that occur during SCI is necessary for the development of promising molecular targets and therapeutic strategies. This study was designed to explore differentially expressed genes (DEGs) associated with the acute and chronic stages of SCI using bioinformatics analysis. Gene expression profiles (GSE45006, GSE93249, and GSE45550) were downloaded from the Gene Expression Omnibus database. SCI-associated DEGs from rat samples were identified, and Gene Ontology and the Kyoto Encyclopedia of Genes and Genomes pathway enrichment analyses were performed. In addition, a protein-protein interaction network was constructed. Approximately 66 DEGs were identified in GSE45550 between 3–14 days after SCI, whereas 2418 DEGs were identified in GSE45006 1–56 days after SCI. Moreover, 1263, 195, and 75 overlapping DEGs were identified between these two expression profiles, 3, 7/8, and 14 days after SCI, respectively. Additionally, 16 overlapping DEGs were obtained in GSE45006 1–14 days after SCI, including \( \textit{Pank1} \), \( \textit{Hn1} \), \( \textit{Tmem150c} \), \( \textit{Rgd1309676} \), \( \textit{Lpl} \), \( \textit{Mdh1} \), \( \textit{Nnt} \), \( \textit{Loc100912219} \), \( \textit{Large1} \), \( \textit{Baiap2} \), \( \textit{Slc24a2} \), \( \textit{Fundc2} \), \( \textit{Mpr14} \), \( \textit{Slc16a7} \), \( \textit{Obf1} \), and \( \textit{Alpk3} \). Importantly, 3882 overlapping DEGs were identified in GSE93249 1–6 months after SCI, including 3316 protein-coding genes and 567 long non-coding RNA genes. A comparative analysis between GSE93249 and GSE45006 resulted in the enrichment of 1135 overlapping DEGs. The significant functions of these 1135 genes were correlated with the response to the immune effector process, the innate immune response, and cytokine production. Moreover, the biological processes and KEGG pathways of the overlapping DEGs were significantly enriched in immune system-related pathways, osteoclast differentiation, the nuclear factor-\( \kappa \)B signaling pathway, and the chemokine signaling pathway. Finally, an analysis of the overlapping DEGs associated with both acute and chronic SCI, assessed using the expression profiles GSE93249 and GSE45006, identified four overlapping DEGs: \( \textit{Slc16a7} \), \( \textit{Alpk3} \), \( \textit{Lpl} \) and \( \textit{Nnt} \). These findings may be useful for revealing the biological processes associated with SCI and the development of targeted intervention strategies.

Key Words: bioinformatics; differential expression; factor; gene; immune response; injury; pathways; protein; spinal cord

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Introduction

Spinal cord injury (SCI) refers to severe trauma to the spinal cord and is characterized by high incidence and morbidity (Shi et al., 2020; Wang et al., 2020). SCI is an overwhelming neurological disorder and has devastating physiological symptoms, which can vary widely from pain, to paralysis, to incontinence (van den Berg et al., 2010; Singh et al., 2014; Jazayeri et al., 2015). Traumatic SCI can be classified into primary injury and secondary injury. Primary injury refers to the mechanical and direct damage incurred against the spinal cord (Stahel et al., 2012), whereas secondary injury refers to the pathological responses induced by the immediate injury. SCI results in complex consequences that involve various changes in biological processes, including the breakdown of the blood-spinal cord barrier, neuroinflammation, oxidative stress, necrosis, apoptosis, glial scar formation, ischemic dysfunction, and functional problems, such as neuropsychiatric pain, autonomic dysfunction, muscle atrophy, and incontinence of the bladder, rectum, and anus (Wyndaele and Wyndaele, 2006; Kottipatruni et al., 2011; Wanner et al., 2013; Anwar et al., 2016).

Despite advances in spinal cord surgical techniques and the efforts devoted to the treatment of SCI and patient care, no effective therapeutic solutions for this destructive neurological disease currently exist (Calvert et al., 2019; Lavis and Goetz, 2019; Thomaz et al., 2019). Owing to the complexity of changes that occur in gene expression and signaling pathways following SCI, understanding the underlying molecular mechanisms associated with pathological events represents a critical step in the development of promising molecular targets and therapeutic strategies. Some studies have attempted to understand this process by identifying differentially expressed genes (DEGs), using microarrays or RNA-sequencing analysis, to reveal the pathways and molecular targets that are activated during acute or chronic SCI (Nesic et al., 2002; van den Berg et al., 2010; Byrnes et al., 2011; Chen et al., 2013; Duran et al., 2017; Du et al., 2018; Guo et al., 2019; Liu et al., 2020). The gene expression profiles GSE45006, GSE93249, and GSE45550 were derived from rat SCI models across different time points after injury and have been used to analyze DEGs and their associated biological processes, based on Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway analyses (Wen et al., 2016; Zhang and Wang, 2016; Duran et al., 2017; Liu et al., 2017; Ma et al., 2017; Zang et al., 2017). Previous studies have focused on temporal gene expression, molecular pathway changes, and changes in the transcriptional regulatory network and molecular mechanisms associated with SCI over time. In addition, comprehensive investigations have examined changes in the expression of both coding and long non-coding genes during the acute and chronic stages of SCI using RNA-sequencing analysis (Duran et al., 2017; Shi et al., 2017; Zhou et al., 2018).

The majority of SCI patients are in the chronic phase of SCI owing to the lack of a cure (Fehlings and Perrin, 2006). However, the pathological changes that are induced during the early stages of SCI might contribute to the development or aggravation of secondary injuries. A major barrier to the identification of ideal therapeutic strategies remains our limited knowledge regarding the coordinated pathophysiological processes induced by SCI. Therefore, determining both the differences and similarities in the molecular changes that characterize acute and chronic SCI is necessary to develop a better understanding of the underlying molecular mechanisms and to develop potential therapeutic strategies. In this study, the expression profiles GSE45006, GSE93249, and GSE45550 were used to perform a comparative analysis of the SCI-related DEGs that change over time and during different phases.

Materials and Methods

Microarray data preprocessing

The gene expression profiles GSE45006, GSE45550, and GSE93249 were downloaded from the Gene Expression Omnibus database of the National Center of Biotechnology Information (http://www.ncbi.nlm.nih.gov/geo/). These expression profiles were based on the GPL1355 platform (GSE45006 and GSE45550) of the Affymetrix Rat Genome 230 2.0 Array and the GPL14844 platform (GSE93249) of Illumina HiSeq 2000. GSE93249 included 11 rat spinal cord samples, including 3 non-injured spinal cord samples in the sham group and 8 SCI samples in the SCI group, representing the chronic phase at 1 month (m1, n = 3), 3 months (m3, n = 3), and 6 months (m6, n = 2) after injury. GSE45550 included 30 rat spinal cord samples, including 6 non-injured spinal cord samples in the sham group and 24 SCI samples representing the acute and subacute phases, at 3 days (d3, n = 8; acute phase), 8 days (d8, n = 8; acute phase), and 14 days (d14, n = 8; subacute phase) after injury. GSE45006 included 24 rat spinal cord samples, including 4 non-injured spinal cord samples in the sham group and 20 SCI samples at 1 day (d1, n = 4; acute phase), 3 days (d3, n = 4; acute phase), 7 days (d7, n = 4; acute phase), 14 days (d14, n = 4; subacute phase), and 56 days (d56, n = 4; chronic phase) post-injury. After the data associated with GSE45006, GSE93249 and GSE45550 were downloaded, the microarray data were pre-processed as follows: the data were converted into expression measures, followed by background correction, quantile normalization, log2 conversion, batch correction, and probe summarization. Finally, the gene expression matrix was obtained.

The identification of DEGs between SCIs and sham samples

The DEGs between SCI and sham samples were screened using GEO2R (http://www.ncbi.nlm.nih.gov/geo/geo2r) online tools (Davis and Meltzer, 2007). The false discovery rate (FDR) was obtained by adjusting the raw P-values with the Benjamini–Hochberg method (Madar and Batista, 2016). The fold-change of gene expression (log, FC) and the FDR were used to select significant DEGs. Genes with log, FC > 1.5 and FDR < 0.05 were selected as significant DEGs.

GO and KEGG pathway enrichment analyses of the DEGs

GO analysis has been widely used to identify functional enrichment patterns in studies examining large-scale genetic sequencing. The KEGG database contains known genes and their associated pathways. The biological significance of DEGs associated with SCI can be assessed by performing GO and KEGG pathway enrichment analyses. The Database for Annotation, Visualization, and Integrated Discovery (DAVID) provides a comprehensive set of functional annotation tools. DAVID was used to identify DEG-associated pathways by calculating P-values (Huang da et al., 2009). The raw P-values were adjusted using the Benjamini-Hochberg method, and FDR < 0.05 was selected as the cut-off criterion.

Statistical analysis

SPSS 22.0 (IBM, Armonk, NY, USA) was used for all statistical analyses. Comparisons among multiple groups were analyzed using a one-way analysis of variance, followed by Dunnett's post hoc test. For experiments involving two groups, an unpaired Student's t-test was performed. A value of P < 0.05 was considered statistically significant. Data are expressed as the mean ± standard deviation (SD).

Results

The DEGs identified between acute SCI and sham groups

The gene expression profile GSE45550 was used to analyze DEGs between the acute SCI and sham groups. Three time points (3, 8, and 14 days after injury) were examined in the experimental groups. The results identified approximately
2411, 463, and 184 DEGs at 3, 8, and 14 days post-injury, respectively. In addition, only 66 overlapping DEGs were identified across the three time points examined between 3 and 14 days post-injury (Figure 1A). The gene expression profile GSE45006 was also examined to further assess DEGs although this profile had a much wider time frame (1, 3, 7, 14, and 56 days post-injury). The results identified approximately 2712 overlapping DEGs during days 1 to 14 post-injury (Figure 1B). Only 2418 DEGs were identified at 56 days post-injury. The above results suggested that the data obtained from GSE45550 and GSE45006 were different.

The overlapping DEGs associated with the acute phase of SCI
Owing to the observed differences between the GSE45550 and GSE45006 profiles, a contrast analysis was performed between these two gene expression profiles for each similar point. More DEGs were identified in GSE45006 than in GSE45550, and approximately 1263, 195, and 75 overlapping DEGs were identified between the two profiles at 3, 7/8, and 14 days post-injury, respectively (Figure 2A–C). The above data indicated that the number of common DEGs significantly decreased with time following injury occurrence. The samples included in GSE45550 were obtained from 3 days to 14 days post-injury, whereas the samples included in GSE45006 covered a much wider time frame (1, 3, 7, 14, and 56 days post-injury). Therefore, GSE45006 was utilized to further determine overlapping DEGs associated with acute SCI over time. As shown in Figure 2D, approximately 16 overlapping DEGs were identified from 1 to 14 days post-injury, including Pank1, Hn1, Tmem150c, Rgd1309676, Lpl (lipoprotein lipase), Mdh1, Nnt (nicotinamide dehydrogenase), Loc100912219, Large1, Baiap2, Slc24a2, Fundc2, Mrps14, Slc16a7 (solute carrier family 16 member 7), Obfc1, and Alpk3 (alpha kinase 3).

The DEGs identified between chronic SCI and sham groups
The gene expression profile GSE93249 was used to screen DEGs between chronic SCI and sham groups. Three time points (1, 3, and 6 months) post-SCI were adopted in the experimental groups. The results indicated approximately 8096, 9683, and 7077 DEGs identified at 1, 3, and 6 months post-injury, respectively. As shown in Figure 3A, approximately 3882 overlapping DEGs were associated with chronic SCI across all time points from 1 to 6 months. Further analysis indicated that these DEGs contained both coding and long non-coding genes, including 3316 protein-coding genes and 567 long non-coding RNAs (Figure 3B and C).

The overlapping DEGs associated with the chronic phase of SCI
To further examine the overlapping DEGs associated with chronic SCI, the expression profiles GSE93249 and GSE45006 were compared. As shown in Figure 4A, a total of 1135 overlapping DEGs were identified between the two profiles when comparing the time points of 1, 3, 6 months, and 56 days (approximately 2 months) post-injury. The 20 most significantly enriched functions associated with the overlapping DEGs, as assessed by GO functional analysis, are shown in the heatmap in Figure 4B. The most striking functions were the immune effector process and the innate immune response. Other significant functions included cytokine production and the positive regulation of cell migration. The enriched biological process and KEGG pathways associated with the overlapping DEGs are shown in Figure 4C. Significantly enriched pathways were associated with the regulation of immune responses, such as the Toll-like receptor (TLR) signaling pathway, osteoclast differentiation, the nuclear factor-(NF)-κB signaling pathway, and the chemokine signaling pathway.

The overlapping DEGs associated with both acute and chronic SCI
To identify common DEGs associated with both acute and chronic phases of SCI, all three expression profiles, GSE45550, GSE93249, and GSE45006, were compared. As shown in Figure 5A and B, only four overlapping DEGs were found in all three gene expression profiles at all time points (acute: 1, 3, 7/8, 14 days; chronic: 1 month, 56 days, 3, 6 months): Slc16a7, Alpk3, Lpl and Nnt. The expression of these four genes, associated with both acute and chronic spinal cord injuries, was further analyzed in GSE93249 and GSE45006 (Figure 6), and the protein-protein interaction networks associated with Slc16a7, Alpk3, Lpl and Nnt were constructed (Additional Figure 1).

Discussion
The pathological changes that occur after traumatic SCI are initially associated with the immediate damage, followed...
by secondary cascade events. Until now, most studies have concentrated on the biomolecular changes and injury-related mechanisms associated with a single phase of SCI: acute, subacute, or chronic. In this study, the time-frame for the experiments was extended from 1 day to 6 months post-injury by analyzing three gene expression profiles, allowing us to conduct a comprehensive overview of SCI.

The events associated with the SCI process can be divided into several contiguous phases, including immediate (0–2 hours), early acute (2–48 hours), subacute (2–14 days), subchronic, and chronic stages (Chen et al., 2013; Duran et al., 2017). This study aimed to provide an overall analysis of common DEGs, with persistent differential expression between the acute (1 day), subacute (7–14 days), and chronic (6 months) phases of SCI. The present study was the first to perform a comparative analysis using the gene expression profiles GSE93249, GSE45550, and GSE45006, which covered a wider time course than has been used in other studies, ranging from 1 day to 6 months post-injury. The three gene expression profiles used in this study were acquired from SCI models based on different injury positions and severity, likely causing differences in the affected signaling pathways and functional molecules. However, we hypothesized that common, shared events were likely to occur, even in different SCI models, the identification of which would provide promising candidates for therapeutic or molecular mechanism studies that may be common among all SCI injuries. DEG clusters were identified, followed by GO and pathway enrichment analyses. The results of this study may provide deep insight into the molecular mechanisms associated with SCI and facilitate the development of novel therapeutic strategies. To ensure the efficiency of DEG screening, only those molecules that were significantly differentially regulated during acute and chronic SCI were selected for our analysis procedures. Comparative analyses revealed that DEGs varied across time points in acute and chronic SCI. Our results identified 16 overlapping DEGs identified across time points in the acute and subacute phases, from 1 to 14 days post-injury. However, when DEGs were examined through 6 months post-injury, only four overlapping DEGs were identified across acute and chronic SCI: Lpl, Nnt, Slc16a7, and Alpk3. The remaining overlapping DEGs identified, in acute but not chronic SCI, were Pank1, Hn1, Tmem150c, Rgd1309676, Mdh1, Loc100912219, Large1, Baiap2, Slc2a4a, Fundc2, Mrps14, and Obfc1. These 12 genes represent acute and subacute SCI-specific genes. These results suggested that early SCI DEGs are distinctly different late-stage SCI DEGs. Thus, these identified genes may represent potential and meaningful candidates for exploring the mechanisms associated with SCI development and the development of strategies to halt or slow SCI progression.

Figure 3 | DEGs between chronic SCI and sham samples.
(A) The gene expression profile GSE93249 was used to analyze DEGs associated with chronic SCI at three time points, (B and C) The overlapping DEGs across 1-, 3-, and 6-month time points after SCI were obtained. Among these DEGs, both (B) coding and (C) long non-coding genes were identified. DEG: Differentially expressed gene; GSE93249: a gene expression profile, which was obtained from the Gene Expression Omnibus database of the National Center of Biotechnology Information; SCI: spinal cord injury.

Figure 4 | The overlapping DEGs associated with chronic SCI across multiple time points.
The expression profiles GSE93249 and GSE45006 were used for comparative analysis to identify overlapping DEGs associated with chronic SCI. (A) The overlapping DEGs were enriched between GSE93249 and GSE45006 from 1 month to 6 months. (B) The significantly enriched functions of the overlapping DEGs are indicated in a heatmap for selected GO analysis. (C and D) The enriched biological processes (C) and KEGG pathways (D) associated with overlapping DEGs. DEG: Differentially expressed gene; GO: gene ontology; GSE93249 and GSE45006: gene expression profiles, which were obtained from the Gene Expression Omnibus database of the National Center of Biotechnology Information; KEGG: Kyoto Encyclopedia of Genes and Genomes; SCI: spinal cord injury.
The inflammatory response induced by injuries to the spinal cord can occur within minutes. Previous studies have identified various immune responses to be the most significantly upregulated biological processes during acute SCI based on GO functional analysis, including the activation of the complement system, the induction of innate and adaptive immune responses, and antibody production (Pineau and Lacroix, 2007; de Rivero Vaccari et al., 2008). Additionally, some chemokines and cytokines produced following SCI have been shown to be involved in the inflammatory responses (Beck et al., 2010; Zhang et al., 2011; Garcia et al., 2016). In this study, one interesting molecule, peroxiredoxin-like 2A (Rgd1309676), was identified during acute and subacute SCI. Peroxiredoxin-like 2A is an antioxidant involved in the redox regulation of the cell, and it functions as a negative regulator of macrophage-mediated inflammation by inhibiting the production of inflammatory cytokines by macrophages, likely through the suppression of the mitogen-activated protein kinase signaling pathway and regulation of osteoclast differentiation (Xu et al., 2010, 2015; Guo et al., 2015).

Therefore, peroxiredoxin-like 2A may function as a negative regulator that combats the excessive inflammation induced by immediate injury during the subacute stage, from days 10 to 14 post-injury. SCI has been shown to be accompanied by a low-grade, chronic inflammatory state, which may be due to the inhibition of excessive inflammation during early stages. Based on the potential function of peroxiredoxin-like 2A and its expression during acute SCI, this molecule may represent a promising and meaningful candidate for exploring SCI-associated cellular mechanisms and therapeutic targets.

Another functional protein, LPL, was identified throughout both the acute and chronic phases of SCI, in this study. LPL has been reported to respond to both SCI-induced inactivity and reloading with training (Baligand et al., 2015). LPL expression has been detected in the brain, spinal cord, and peripheral nerves. In addition to the hydrolysis of triglyceride-rich lipoproteins, LPL has been shown to play important roles in the pathophysiologic responses of nerve injury. Several studies have reported the selective induction of LPL expression in response to cerebral ischemia-reperfusion injury and have suggested a protective function of LPL, which facilitates lipid uptake and reutilization in the central nervous system during demyelination events or following nerve crush injuries (Blain et al., 2004; Paradis et al., 2004; Wang et al., 2010; Bruce et al., 2018). Based on our results, LPL may fulfill biological functions during the pathophysiologic responses to both acute and chronic SCI. Therefore, LPL may represent a protective factor, playing potential roles in the clearance of myelin-derived lipids after SCI.

NNT is essential for the removal of hydrogen peroxide (Meimari et al., 2018), and currently studies examining NNT primarily focus on adrenal diseases, heart diseases, diabetes, and other diseases (Chatterjee et al., 2014; Hershkovitz et al., 2015; Kampjut and Sazanov, 2019).

SLC16A7 is a member of the monocarboxylate transporter family, and has been reported to be expressed at increased levels in prostate and colorectal cancer specimens compared with control specimens (Pinheiro et al., 2008; Pertega-Gomes et al., 2015). ALPK3 plays a key role in transerase activity, the transfer of phosphorus-containing groups, and protein serine/threonine kinase activity (Feng and Jiang, 2020). Although these three genes were identified during the acute and chronic stages of SCI, relatively few reported studies have examined these genes in nerves and the spinal cord, so their significance to the nervous system remains unclear.

In summary, during the present study, we performed an overall bioinformatics analysis of DEGs associated with acute, subacute, and chronic SCI through the utilization and analysis of three gene expression profiles. Approximately 120 DEGs and 1135 DEGs were associated with the acute phase and the chronic phase of SCI, respectively. However, only four overlapping DEGs, Slc16a7, Alpk3, Lpl, and Nnt, displayed persistent differential expression across the acute, subacute, and chronic phases of SCI. Thus, the twelve DEGs that were...
associated specifically with the acute phases of SCI may be associated with early pathological changes that occur after SCI. In contrast, the four shared genes identified in all phases of SCI may function to exert constant biological effects during SCI progression. Taken together, these data might provide some meaningful insight regarding differences in the pathogenic mechanisms and targeted therapy strategies between acute and chronic SCI. However, because these are predictive results and are based on bioinformatics analyses, the functions of key DEGs, such as peroxiredoxin-like 2A and LPL, and their roles in SCI remain to be explored and demonstrated by biological experiments. SCI models associated with different injury positions and severity are likely to result in varying DEG expression; therefore, the DEGs identified in this study may be further verified in different models in future studies.

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Additional file: Additional Figure 1: The protein-protein interaction network.

References

Anwar MA, Al Shehahi TS, Eid AH (2016) Inflammamgenesis of secondary spinal cord injury. Front Cell Neurosci 10:98.

Baglad C, Chen YW, Ne P, Pandey SN, Liu SH, Liu M, Vandemark RE (2015) Transcriptational pathways associated with skeletal muscle changes after spinal cord injury and treadmill locomotor training. Biomed Res Int 2015:387090.

Beck KD, Nguyen HX, Galvan MD, Salo KD, Woodruff TM, Anderson AJ (2010) Quantitative analysis of cellular inflammation after traumatic spinal cord injury: evidence for a multiphasic inflammatory response in the acute to chronic environment. Brain 133:433-447.

Blain IF, Paradies C, Gaudreault SB, Champagne D, Richard D, Poirier J (2018) Lipoprotein lipase (LPL) is a Nature of alternatively-activated microglia and may facilitate lipid uptake in the CNS during demyelination. Front Mol Neurosci 11:57.

Calvert S, Grunin PJ, Zhao KD, Lee KH (2019) Emergence of epidural electrical stimulation to facilitate sensorimotor reorganization functionally after spinal cord injury. Neuroplast Modulat Neuroregul 2019:244-252.

Chatterjee S, Chakraborty A, Weinberg I, Kadakia M, Wilensky RL, Sardar P, Kumbhani DJ, Mukherjee D, Byrnes KR, Washington PM, Knoblach SM, Hoffman E, Faden AI (2011) Delayed inflammatory mRNA and protein expression in the acute to chronic environment. Brain 133:433-447.

Bruce KD, Nguyen HX, Galvan MD, Boyle KE, Macklin WB, Eckel RH (2018) Lipoprotein lipase (LPL) is a Nature of alternatively-activated microglia and may facilitate lipid uptake in the CNS during demyelination. Front Mol Neurosci 11:57.

Byrnes KR, Washington PM, Knoblach SM, Hoffman E, Faden AI (2011) Delayed inflammatory mRNA and protein expression in the acute to chronic environment. Brain 133:433-447.

Cho YJ, Wu JQ, Yin Q, Zhang JS, Li L, Wang X, Zou Y, Wang X, Wang Y, Liu L, Xia J, Wang X, Xu X, Wu E (2010) The expression and activity of brain lipoprotein lipase is increased after acute cerebral ischemia-reperfusion in rats. Neuropathology 30:131-139.

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Additional Figure 1 The protein-protein interaction related to Lpl, Nnt, Slc16a7 and Alpk3 were constructed respectively to build a potential interaction network.

Alpk3: Alpha kinase 3; LPL: lipoprotein lipase; Nnt: nicotinamide tide transhydrogenase; Slc16a7: solute carrier family 16 member 7.