6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase Suppresses Neuronal Apoptosis by Increasing Glycolysis and "cyclin-dependent kinase 1-Mediated Phosphorylation of p27 After Traumatic Spinal Cord Injury in Rats

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Abstract
Apoptosis is a vital pathological factor that accounts for the poor prognosis of traumatic spinal cord injury (t-SCI). The 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase (PFKFB3) is a critical regulator for energy metabolism and proven to have antiapoptotic effects. This study aimed to investigate the neuroprotective role of PFKFB3 in t-SCI. A compressive clip was introduced to establish the t-SCI model. Herein, we identified that PFKFB3 was extensively distributed in neurons, and PFKFB3 levels significantly increased and peaked 24 h after t-SCI. Additionally, knockdown of PFKFB3 inhibited glycolysis, accompanied by aggravated neuronal apoptosis and white matter injury, while pharmacological activation of PFKFB3 with meclizine significantly enhanced glycolysis, attenuated t-SCI-induced spinal cord injury, and alleviated neurological impairment. The PFKFB3 agonist, meclizine, activated cyclin-dependent kinase 1 (CDK1) and promoted the phosphorylation of p27, ultimately suppressing neuronal apoptosis. However, the neuroprotective effects of meclizine against t-SCI were abolished by the CDK1 antagonist, RO3306. In summary, our data demonstrated that PFKFB3 contributes robust neuroprotection against t-SCI by enhancing glycolysis and modulating CDK1-related antiapoptotic signals. Moreover, targeting PFKFB3 may be a novel and promising therapeutic strategy for t-SCI.

Keywords
PFKFB3, meclizine, neuronal apoptosis, glycolysis, traumatic spinal cord injury

Introduction
Traumatic spinal cord injury (t-SCI) is one of the most serious injuries worldwide, which can lead to a series of physiological dysfunctions, including motor and sensory impairment, chronic pain, respiratory and cardiovascular alterations, neurogenic bowel, neurogenic bladder, and even psychological disorders¹. The pathogenesis of t-SCI is complicated; it consists of primary trauma followed by secondary damage, further aggravating the injury. The primary injury quickly causes acute, mechanical destruction to the spinal cord tissues, including axons, blood vessels, and cells at the injury site¹. The secondary injury is initiated after the primary insult and progresses throughout the entire
period of injury. Various mechanisms are associated with the secondary injury, including hemorrhage, ischemia, oxidative stress, inflammation, death of neurons and glia, axonal demyelination and degeneration, extracellular matrix remodeling, and gliosis. Additionally, neuronal apoptosis plays a critical role in contributing to the behavioral dysfunction. As a result, multiple drugs aiming to reduce neuronal apoptosis may contribute to improved neurological outcomes following t-SCI. However, the underlying mechanism of neuronal apoptosis has not yet been fully elucidated, and the curative effects are still poor.

The enzyme 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase (PFKFB) is a bifunctional enzyme that is involved in the regulation of glucose metabolism. The PFKFB family contains four types of isozymes, PFKFB1-4. PFKFB3 is the most studied isof orm and can catalyze the synthesis of the allosteric activator, fructose-2,6-bisphosphate (F2,6BP). Increased F2,6BP promotes glycolytic flux by activating the key glycolytic enzyme, phosphofructokinase 1 (PFK1), which is important for lactate production, whereas decreased F2,6BP levels inhibit glycolysis. As a result, PFKFB3 is considered a critical stimulator of glycolysis. Previous studies have suggested that there is enhanced expression of PFKFB3 in rapidly proliferating transformed cells, solid tumors, and leukemias. Cancer cells preferentially utilize glycolysis, rather than the citric acid cycle, as their source of energy, and this is known as the Warburg effect. This allows the tumor cells to quickly utilize glucose, providing energy and materials for enhanced tumor proliferation or the prevention of apoptosis. As a result, PFKFB3 exerts critical antiapoptotic effects toward cancer cells. PFKFB3 is also expressed on mitochondria, as their source of energy, and this is known as the Warburg effect. This allows the tumor cells to quickly utilize glucose, providing energy and materials for enhanced tumor proliferation or the prevention of apoptosis.

Materials and Methods

Animals

Male Sprague Dawley rats, each weighing 250–300 g, used in this study were purchased from SLAC Laboratory Animal Co. Ltd (Shanghai, China). The animals were maintained in an environment under a constant temperature (22 ± 1°C) and humidity (60 ± 5%) with a 12-h light/dark cycle.

t-SCI Model

A spinal cord compressive model of t-SCI was recreated, according to the previous description. In brief, the T10 vertebrae of the rats were exposed, and a laminectomy was conducted to reveal the spinal cord. A vascular clip (30 g force, INS 14120, Kent Scientific, Torrington, CT, USA) was then used to clamp the spinal cord for 30 s to create a compressive injury. The rats in the sham group received the same operation but without the clamp.

Drug and Small Interfering RNA Administration

Meclizine (100 mg/kg, Macklin, Shanghai, China), dissolved in 5 ml of normal saline, was administered to the rats via intraperitoneal injection 1 h after t-SCI. A mixture, consisting of two types of specific small interfering RNA (siRNA) to target the rat PFKFB3 mRNA (Genomeditech, Shanghai, China) at a dosage of 500 pmol, was prepared and dissolved in 10 μl Entranster™ in vivo transfection reagent (Engreen Biosystem, Beijing, China), and then intrathecally injected into the rats 48 h before t-SCI. The scramble siRNA was administered to the rats following the same process. RO3306 (4 mg/kg, Selleck Chemicals, Houston, TX, USA) was administered to the rats via oral gavage 1 h after t-SCI.

Intrathecal Injection

Intrathecal injections were conducted using Hylden’s method, according to the previous description. In brief, a
rat was held in one hand with the back bulged, and a lumbar puncture was conducted using a microsyringe to penetrate the intervertebral space between the L5 and L6 vertebrae. A total of 10 µl of the drug was injected at a rate of 2 µl/min. The syringe was securely positioned for 10 min and then removed. The sham rats underwent the same procedure without drug administration.

**Experimental Designs**

**Experiment 1.** Rats were randomly divided into seven groups: sham, t-SCI 3 h, t-SCI 6 h, t-SCI 12 h, t-SCI 24 h, t-SCI 48 h, and t-SCI 72 h. Western blotting was performed in each group, while immunofluorescence staining was performed in the sham and t-SCI 24 h groups.

**Experiment 2.** Rats were randomly divided into six groups: sham, t-SCI + vehicle, t-SCI + scramble siRNA, t-SCI + PFKFB3 siRNA, t-SCI + meclizine, and t-SCI + meclizine + PFKFB3 siRNA. Then, 24 h after t-SCI, western blotting was performed. The F2,6-BP and lactic acid levels were measured in each group.

**Experiment 3.** Rats were randomly assigned into three groups: sham, t-SCI + vehicle, and t-SCI + meclizine. Basso, Beattie, and Bresnahan (BBB) and combined behavioral scores (CBS) were evaluated before treatment and on the 1st, 3rd, 7th, and 14th day, as well as every 2 weeks following treatment until the eighth week.

**Experiment 4.** Rats were randomly assigned to five groups: sham, t-SCI + vehicle, t-SCI + meclizine, t-SCI + RO3306, and t-SCI + meclizine + RO3306. Then, 24 h after t-SCI, western blotting, immunofluorescence staining, and transmission electron microscopy (TEM) were performed. CDK1 activity and nuclear F2,6-BP were also tested in each group.

Detailed experimental designs are shown in Fig. 1A.

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**Fig. 1.** Experimental designs (A). The time course of PFKFB3, and F2,6BP showed that the levels of PFKFB3 and F2,6BP were significantly elevated at 6 h, peaked at 24 h, and then significantly decreased after 24 h postinjury. Representative western blot images (B); the levels of PFKFB3 (C); and the levels of F2,6BP (D). The expression of FBKFB3 in neurons was increased at 24 h postinjury. Representative microphotographs of double immunofluorescence of PFKFB3 and NeuN (E): the proportion of PFKFB3-positive neurons (F). N = 6 for each group. Data are expressed as the mean ± SD. *P < 0.05 versus sham; #P < 0.05 versus t-SCI 24 h. F2,6BP: fructose-2,6-bisphosphate; PFKFB3: 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase; SD: standard deviation.
Motor Function Assessment
The neurologic functions were evaluated by two well-trained researchers who were blinded to the groups and used the BBB scores and CBS, as previously reported37.

Western Blotting
A spinal cord segment, 0.5 cm in length, from where the injured site was centered was used for protein quantification using western blotting. Proteins of the nucleus and cytosol were extracted using the NE-PER Nuclear and Cytoplasmic Extractions Kit (Thermo Fisher Scientific, Waltham, MA, USA), following the manufacturer’s instructions. The protein mixture extracted from each segment was separated via electrophoresis and transferred to the polyvinylidene difluoride membrane (Bio-Rad Laboratories, Hercules, CA, USA). The membranes were first incubated with PFKFB3 (1:5,000, ab181861, Abcam, Cambridge, UK), Bax (1:1,000, ab32503, Abcam), Bcl-2 (1:500, ab59348, Abcam), cleaved caspase-3 (CC-3, 1:500, ab13847, Abcam), p-CDK1(1:1,000, ab201008, Abcam), p27 (1:5,000, ab32034, Abcam), H3 (1:5,000, ab1791, Abcam), myelin basic protein (MBP) (1:1,000, ab209328, Abcam), APP (1:1,000, ab59348, Abcam), H3 (1:5,000, ab1791, Abcam), myelin basic protein (MBP) (1:1,000, ab209328, Abcam), APP (1:1,000, ab59348, Abcam), and p-p27 (1:1,000, ab75908, Abcam) antibodies at 4°C overnight, respectively, and incubated with the secondary antibodies (1:10,000, ZB-2301 or ZB-2305, Zhongshan Golden Bridge, Beijing, China) at 25°C for 2 h. Finally, 4',6-diamidino-2-phenylindole (1 μg/ml, Roche Inc., Basel, Switzerland) was each incubated at 25°C for 2 h. Finally, 4',6-diamidino-2-phenylindole (1 μg/ml, Roche Inc.) was used to dye the nucleus and mount the medium. A fluorescence microscope (Olympus, Tokyo, Japan) was used to observe the tissue sections, and Photoshop 13.0 software (Adobe Systems Inc., San Jose, CA, USA) was used for photo postprocessing. Six sections were obtained from each sample, and one random field of gray matter was used from each section to count the cell numbers at ×200 magnification.

PFKFB3 expression was conveyed as the mean ratio of PFKFB3-positive neurons to total neurons in each group. Neuronal apoptosis was expressed as the mean ratio of CC-3-/TUNEL-positive neurons to total neurons in each group.

Measurement of Lactic Acid Levels
The detection of tissue lactic acid content was conducted using the Lactic Acid Assay Kit (JianCheng Bioengineering Institute, Nanjing, Jiangsu, China), following the manufacturer’s instructions. The fresh tissues were homogenized in 9 × volume normal saline and centrifuged at 2,500 × g for 10 min at 4°C. Before conducting lactic acid detection, the supernatant was deproteinized via the trichloroacetic acid (TCA) method using the Deproteinizing Sample Preparation Kit II (Biovision, San Francisco, CA, USA). In brief, a sample of 100 μl was mixed with 15 μl of cold TCA, maintained on ice for 15 min, and centrifuged at 12,000 × g for 15 min at 4°C. The supernatant was mixed with 10 μl of cold neutralization solution and maintained on ice for 5 min. A 20 μl sample, 1,000 μl working solution, and 200 μl chromogenic reagent were mixed and incubated at 37°C for 10 min. Then, the terminating solution was added to stop the reaction. The absorbance was measured at 530 nm. The results were expressed as mmol of lactic acid per gram of protein. The data were compared with the sham group to acquire the relative lactic acid levels. The assays were repeated three times independently.

Nuclei Isolation
Nuclei were isolated using Nuclear Extraction Kit (Solarbio, Beijing, China), following the manufacturer’s instructions. In brief, the fresh tissues were mixed with 1 ml of the lysis buffer and 50 μl of reagent A and placed into a homogenizer to grind into a tissue homogenate. The suspension was centrifuged at 700 × g for 5 min at 4°C, and the precipitate was resuspended in 0.5 ml cold lysis buffer. The suspension was then added to 0.5 ml medium buffer in an eppendorf (EP) tube and centrifuged at 700 × g for 5 min at 4°C. The precipitate was resuspended in 0.5 ml cold lysis buffer again and centrifuged at 1,000 × g for 10 min at 4°C to acquire pure nuclei40.

Measurement of F2,6BP Levels in Cells and Nuclei
The F2,6BP content was determined as previously described41. Briefly, tissue debris or pure nuclei were lysed in 0.1 M NaOH and incubated at 80°C for 5 min. After centrifugation, the supernatant was used for the F2,6BP assay after being neutralized with ice-cold acetic acid in 20 mmol/l 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid and maintained on ice for 5 min. The supernatant was mixed with 10 μl of cold neutralization solution and maintained on ice for 5 min. A 20 μl sample, 1,000 μl working solution, and 200 μl chromogenic reagent were mixed and incubated at 37°C for 10 min. Then, the terminating solution was added to stop the reaction. The absorbance was measured at 530 nm. The results were expressed as mmol of lactic acid per gram of protein. The data were compared with the sham group to acquire the relative lactic acid levels. The assays were repeated three times independently.
acid buffer. The supernatant was added into the mixture containing 50 mmol/l Tris/HCl (pH 8.0), 5 mmol/l MgCl₂, 0.15 mmol/l NADH, 17.5 mmol/l glucose-6-phosphate, 0.5 mmol/l pyrophosphate, 1 mmol/l triosephosphate isomerase, and 10 µg/ml aldolase, 1 µg/ml triosephosphate isomerase, and 10 µg/ml glycerol-3-phosphate dehydrogenase and was incubated at 25°C for 2 min. The absorbance was measured at 340 nm. The results were expressed as pmol of F₂,6BP per microgram of protein. The data were compared with those of the sham group to acquire the relative F₂,6BP levels, and the assays were repeated three times independently.

Detection of CDK1 Activity

CDK1 activity was detected using the MESACUP Cdc2/Cdk1 kinase assay kit (MBL, Nagoya, Japan), according to the manufacturer’s instructions. Briefly, the pure nuclei were lysed in 50 µl of hypotonic lysis buffer on ice for 1 h. After centrifugation, 2.5 µl of supernatant, 2.5 µl of 10× cdc2 reaction buffers, 2.5 µl of biotinylated MV peptide, 15 µl of distilled water, and 2.5 µl of 1 mmol/l adenosine triphosphate (ATP) were mixed together and incubated at 30°C for 30 min. Then, 100 µl of phosphorylation terminating reagent was added. After centrifugation, 50 µl of supernatant was added to the wells of a microplate coated with phosphor-MV peptide antibody, which was then incubated at 25°C for 60 min. After washing, 50 µl of peroxidase (POD)–streptavidin conjugate was added to each well, which was then incubated at 25°C for 30 min. After washing again, 50 µl of substrate solution was added to each well, which was incubated at 25°C for 5 min before the terminating reagent was added. The absorbance was measured at 492 nm. The data were compared with those of the sham group to acquire the relative optical density (OD) value. The assays were repeated three times independently.

Fig. 2. Knockdown of PFKFB3 decreased F₂,6BP and lactic acid, while increased apoptosis and white matter injury at 24 h postinjury. Representative western blot images (A); the levels of PFKFB3 (B); the levels of F₂,6BP (C); the Bcl-2/Bax ratio (D); the levels of CC-3 (E); the lactic acid levels (F); the levels of MBP (G); the levels of APP (H). N = 6 for each group. Data are expressed as the mean ± SD. *p < 0.05 versus sham; #p < 0.05 versus t-SCI + vehicle; @p < 0.05 versus t-SCI + scramble siRNA. F₂,6BP: fructose-2,6-bisphosphate; PFKFB3: 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase; SD: standard deviation; siRNA: small interfering RNA; t-SCI: traumatic spinal cord injury.
Transmission Electron Microscopy

A spinal cord segment, 0.5 cm in length, from where the injured site was centered was obtained. The gray matter from the proximal and distal spinal cord (2 mm away from the center of the injury site) were cut into 1 mm³ tissue blocks and fixed overnight in 2.5% glutaraldehyde. Next, the blocks were rinsed, fixed in 1% osmic acid for 1 h, dehydrated in gradient alcohol, and embedded in araldite overnight at 60°C. Finally, the blocks were sectioned into 100 nm ultrathin slices, stained using uranyl acetate and lead citrate, and were then observed using a TEM (Philips Tecnai 10, Amsterdam, Holland). Six sections were obtained from each sample, and 10 random neurons per section were used to count the mitochondria around the neuron nucleus at ×4,200 magnification. The mitochondria vacuolization rate refers to the mean ratio of vacuolated to total mitochondria in each group.

Statistical Analysis

The results were presented as mean ± standard deviation and analyzed by t-test, one- or two-way analysis of variance (ANOVA), and Bonferroni’s post hoc multiple comparisons test, using P < 0.05 as the standard for statistical significance. Statistical analyses were conducted using GraphPad Prism 6 (San Diego, CA, USA).
Fig. 4. Meclizine improved motor function of t-SCI rats. The BBB scores (A); the CBS (B). N = 6 for each group. Data are expressed as the mean ± SD. *P < 0.05 versus sham; #P < 0.05 versus t-SCI + vehicle. BBB: Basso, Beattie, and Bresnahan; dpi: day postinjury; SD: standard deviation; t-SCI: traumatic spinal cord injury.

Fig. 5. Meclizine’s antiapoptotic effects were mediated by CDK1 and could be reversed by RO3306 at 24 h postinjury. Representative western blot images (A); the levels of nuc-PFKFB3 (B); the levels of nuc-F2,6BP (C); the CDK1 activities (D); the levels of nuc-p-CDK1 (E); the nuc-p27/nuc-p27 ratio (F); the Bcl-2/Bax ratio (G); and the levels of CC-3 (H). N = 6 for each group. Data are expressed as the mean ± SD. *P < 0.05 versus t-SCI + vehicle; #P < 0.05 versus t-SCI + meclizine; @P < 0.05 versus t-SCI + RO3306. CDK1: cyclin-dependent kinase 1; F2,6BP: fructose-2,6-bisphosphate; PFKFB3: 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase; SD: standard deviation; t-SCI: traumatic spinal cord injury.
Results

Time Course of FBKFB3 and F2,6BP

PKFB3 level began increasing significantly at 6 h following t-SCI, reaching its peak at 24 h, compared to the sham group ($P < 0.05$). The PFKFB3 level then gradually decreased at 48 and 72 h following t-SCI, compared to the t-SCI 24 h group ($P < 0.05$, Fig. 1B, C). Interestingly, the F2,6BP levels exhibited similar trends (Fig. 1D).

Expression of FBKFB3 in Neurons 24 h After t-SCI

PKFB3 was expressed in both the nucleus and cytoplasm of neurons. The t-SCI 24 h group had a higher proportion of PFKFB3-positive neurons than the sham group ($P < 0.05$, Fig. 1E, F).

Knockdown of PFKFB3 Decreased Lactic Acid Levels and Increased Apoptosis and White Matter Injury 24 h After t-SCI

When compared to the sham group ($P < 0.05$), the t-SCI + vehicle group had significantly elevated levels of PFKFB3, F2,6BP, lactic acid, CC-3, and APP, but the Bcl-2/Bax ratio and the level of MBP were significantly decreased. PFKFB3 siRNA significantly decreased the PFKFB3, F2,6BP, lactic acid, and MBP levels, as well as the Bcl-2/Bax ratio, but increased the levels of CC-3 and APP, compared to the t-SCI + vehicle group ($P < 0.05$). In addition, the scramble siRNA did not alter the PFKFB3 levels, nor its downstream molecules compared to the t-SCI + vehicle group ($P > 0.05$, Fig. 2A–H).

Upregulation of PFKFB3 Increased Lactic Acid and Reduced Apoptosis and White Matter Injury 24 h After t-SCI

Meclizine significantly increased the PFKFB3, F2,6BP, lactic acid, and MBP levels, as well as the Bcl-2/Bax ratio, but decreased the levels of CC-3 and APP, compared to the t-SCI + vehicle group ($P < 0.05$). Moreover, the combined treatment with PFKFB3 siRNA significantly reversed the effects of meclizine ($P < 0.05$ t-SCI + meclizine versus t-SCI + meclizine + PFKFB3 siRNA, Fig. 3A–H).

Meclizine Improved Motor Function

The BBB scores slightly decreased on the first day but rapidly returned to near-baseline levels in the sham group.
However, the BBB scores in the t-SCI + vehicle group and the t-SCI + meclizine group decreased to almost 0 on the first day after t-SCI, which were significantly lower than those in the sham group ($P < 0.05$). As observations continued, the BBB scores gradually increased in these two groups but were still significantly lower than those in the sham group ($P < 0.05$). On the 14th day and subsequent periods of observation, the BBB scores in the t-SCI + meclizine group showed a marked increase compared to those in the t-SCI + vehicle group ($P < 0.05$, Fig. 4A).

The CBS of the sham group slightly increased on the first day, but rapidly returned to a score near 0. However, the CBS in both the t-SCI + vehicle group and the t-SCI + meclizine group increased to a score of approximately 100 on the first day after t-SCI, which was noticeably higher than the scores in the sham group ($P < 0.05$). As observations continued, CBS gradually decreased in these two groups but were still evidently higher than those in the sham group ($P < 0.05$). Meclizine significantly increased CBS on the 14th day and the subsequent observation points when compared to the t-SCI + vehicle group ($P < 0.05$, Fig. 4B).

The proportions of CC-3- and TUNEL-positive neurons significantly increased in the t-SCI + meclizine group compared to the t-SCI + vehicle group ($P < 0.05$). The combined treatment with RO3306 reversed these effects on the 14th day and the subsequent observation points when compared to the t-SCI + vehicle group ($P < 0.05$, Fig. 4B).

**Role of CDK1 in the Meclizine-mediated Antiapoptotic Effects 24 h After t-SCI**

Meclizine significantly increased the levels of nuc-PFKFB3 and nuc-F2,6BP compared to the t-SCI + vehicle group ($P < 0.05$). The combined treatment with RO3306 did not alter the levels of nuc-PFKFB3 and nuc-F2,6BP ($P > 0.05$). The CDK1 activity, the level of nuc-p-CDK1, the nuc-p-p27/nuc-p27 ratio, and the Bcl-2/Bax ratio significantly increased, and the level of CC-3 significantly decreased in the t-SCI + meclizine group compared to the t-SCI + vehicle group ($P < 0.05$). However, combined treatment with RO3306 reversed these effects ($P < 0.05$ t-SCI + meclizine + RO3306 versus t-SCI + meclizine, Fig. 5A–H).

The proportions of CC-3- and TUNEL-positive neurons significantly increased in the t-SCI + vehicle group as compared to the sham group ($P < 0.05$). Meclizine had the opposite effect and significantly decreased the proportions of CC-3- and TUNEL-positive neurons compared to the t-SCI + vehicle group ($P < 0.05$). The combined treatment with RO3306 reversed these effects ($P < 0.05$ t-SCI + meclizine + RO3306 versus t-SCI + meclizine, Figs. 6 and 7).

Electron microscopy showed that the nuclei and mitochondria in the neurons of the sham group appeared normal. (Fig. 7.) Meclizine’s antiapoptotic effects were reversed by RO3306 at 24 h postinjury. Representative microphotographs of double immunofluorescence staining of TUNEL and NeuN (A); the proportion of TUNEL-positive neurons (B). $N = 6$ for each group. Data are expressed as mean ± SD. $^aP < 0.05$ versus t-SCI + vehicle; $^bP < 0.05$ versus t-SCI + meclizine; $^{*}P < 0.05$ versus t-SCI + RO3306. SD: standard deviation; t-SCI: traumatic spinal cord injury; TUNEL: terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling.
However, the neurons in the t-SCI + vehicle group displayed abnormal ultrastructure, including karyopyknosis, chromatin condensation, and margination, as well as swelling, vacuolization, and reduction of mitochondria. Meclizine recovered the normal ultrastructure, while combined treatment with RO3306 reversed the effects of meclizine. In the t-SCI + vehicle group, the proportions of mitochondrial vacuolization significantly increased when compared to the sham group ($P < 0.05$) and decreased by meclizine ($P < 0.05$). Combined treatment with RO3306 significantly reversed the effects of meclizine ($P < 0.05$, Fig. 8A, B).

**Discussion**

In this study, we investigated the role of PFKFB3 in the context of t-SCI and further explored the underlying mechanism. The major findings were: (1) the level of neuronal PFKFB3 significantly increased and peaked at 24 h after t-SCI; (2) knockdown of PFKFB3 inhibited glycolysis and aggravated neuronal apoptosis and white matter injury, while pharmacological activation of PFKFB3 with meclizine significantly enhanced glycolysis and attenuated t-SCI and neurological impairment; (3) meclizine activated CDK1 and promoted the phosphorylation of p27; and (4) pharmacological inhibition of CDK1 reversed the meclizine-mediated antiapoptotic effect under t-SCI conditions. Based on this evidence, PFKFB3 was found to contribute robust neuroprotection against t-SCI by enhancing glycolysis and modulating CDK1-related antiapoptotic signals (Fig. 9).

Apoptosis, characterized by distinctive morphologic changes, such as cell shrinkage, nuclear fragmentation, chromatin condensation, and chromosomal DNA fragmentation, is a programmed cell death process that has been proven to...
be a critical pathological mechanism accounting for the poor outcome of multiple human diseases, such as heart failure, pulmonary fibrosis, and liver injury. Additionally, recent studies have demonstrated that targeting apoptosis is beneficial for several acute and chronic CNS diseases, including cerebral ischemia, intracerebral hemorrhage, and Alzheimer’s disease. Despite the great efforts that have been made, the exact mechanism of apoptosis has not yet been fully elucidated. Notably, recent studies have revealed an important role of PFKFB3, a critical regulator for glycolysis and metabolism, in the pathological process of apoptosis. The PFKFB3 protein is expressed extensively in human tissues, such as the heart, kidneys, intestines, and spinal cord. The activated PFKFB3 can upregulate the activity of F2,6BP, which further activates PFK1, resulting in enhanced glycolysis. Increased glycolysis may confer a beneficial effect for cellular survival via multiple mechanisms, including increasing energy supplements, inhibiting the production of reactive oxygen species, and maintaining the biofunction of voltage-dependent anion channels. Notably, the upregulation of glycolysis has been proven to exert a robust neuroprotective effect in ischemic stroke and Alzheimer disease (AD). Despite the beneficial effect of PFKFB3 being studied for years, little is known regarding its role in the pathological process of t-SCI.

Therefore, in the first part of this study, we explored the temporal pattern of PFKFB3 after t-SCI. Additionally, similar to a previous study reporting an upregulated level of PFKFB3 after endotoxemia-induced myocardial injury, we observed significant activation of PFKFB3, evidenced by increased PFKFB3 expression and upregulated levels of its substrate F2,6BP, after t-SCI. We then further investigated the effect of PFKFB3 under t-SCI conditions. We found that, consistent with a previous study, inhibition of PFKFB3 significantly inhibited glycolysis, as reported, and meclizine exerted significant neuroprotective effects by attenuating t-SCI-induced neuronal apoptosis and white matter injury. This suggests a critical role of PFKFB3 in regulating glycolysis-related apoptosis under t-SCI conditions, which is a similar result to that which has previously been reported.

Notably, other than regulating glycolysis, recent studies have revealed that PFKFB3 can exert antiapoptotic effects via a glycolysis-independent pattern. The increased F2,6BP levels, which were induced by PFKFB3, could translocate...
into nuclei. Therefore, the nuclear F2,6BP could activate CDK1 and subsequently promote the phosphorylation and ubiquitination degradation of p27, a critical initiator to apoptosis, leading to the prevention of cellular apoptosis. Multiple molecular events may cause the enhancement of CDK1 activity, thus, promoting the cell cycle, while the inhibition of CDK1 leads to cell cycle arrest and apoptosis. Further study of F2,6BP may be a novel and promising therapeutic strategy related antiapoptotic signals. Our data support that targeting F2,6BP mediated by enhancing glycolysis and modulating CDK1-related white matter injury. The mechanisms are possibly against t-SCI by suppressing neuronal apoptosis and reducing white matter injury. The mechanisms are possibly against t-SCI by suppressing neuronal apoptosis and reducing white matter injury.

Conclusions

Our data support that the pharmacological activation of PFKFB3 with meclizine conferred a robust beneficial effect against t-SCI by suppressing neuronal apoptosis and reducing white matter injury. The mechanisms are possibly mediated by enhancing glycolysis and modulating CDK1-related antiapoptotic signals. Our data support that targeting PFKFB3 may be a novel and promising therapeutic strategy for t-SCI.

Limitations

There were some limitations in our study. First, this study only focused on antiapoptotic effects of PFKFB3 without investigation of its other roles, such as autophagy, insulin signaling, and inflammatory roles, which should be further explored. Second, the antiapoptotic pathway of PFKFB3 in this study was limited to glycolysis enhancement and CDK1-related p27 phosphorylation; further studies on the relationship of PFKFB3 and other signal pathways in neuronal apoptosis after t-SCI are also required.

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