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Abstract

Electro acupuncture has been widely used to treat demyelinating diseases, such as multiple sclerosis (MS), acute haemorrhagic leukoencephalitis (AHLE), compressed spinal cord injury (CSCI). However, the protective effect of electro acupuncture (EA) on neural myelin sheaths remains controversy attributed to the dimness of its therapeutic mechanism. In this study, we tried to explore the protective mechanism of EA in a custom-designed model of CSCI. Zusanli (ST36) and Huantiao (GB30) acupoints were stimulated by EA. The motor functions were monitored by Basso, Beattie and Bresnahan locomotor rating scale. The pathological changes in axonal myelinated fibers were estimated by luxol fast blue (LFB) and transmission electron microscopy (TEM). Epidermal growth factor factor receptor (EGFR), oligodendrocyte transcription factor 2 (Olig2), caspase-3 and phosphorylated Akt1 (pAkt1) were detected by immunofluorescence and western blot assays. After 7-day treatment of EA, the expression of p-EGFR, pAkt1 and Olig2 was significantly up-regulated which was consistent with changes of locomotor skill and ultrastructure of myelin sheath. By contrast, the expression of active caspase-3 was obviously down-regulated. Our results indicated that the protective effect of EA on neural myelin sheaths might be mediated via activation of EGFR after CSCI.

Keywords: Electro-acupuncture; Compressed spinal cord injury; Remyelination; p-EGFR; pAkt1; Olig2; Caspase-3

Introduction

Electro acupuncture is a form of acupuncture where a small electric current is passed between pairs of acupuncture needles. It can not only augment the use of regular acupuncture, but also introduce electric current into human body. By promoting oligodendrocyte proliferation and differentiation, electro acupuncture (EA) has been widely used in demyelinating diseases and showed excellent curative effect [1-4]. However, the mechanism of EA is largely unknown.

Compressed spinal cord injury (CSCI), which is caused by a metastatic extradural tumor, lumbar intervertebral disc herniation, has become a global issue. Previous studies have proved that extensive demyelination induced by oligodendrocytes apoptosis was the major pathological change after CSCI [5-7], suggesting that the main goal in treatment of CSCI is the protection of oligodendrocytes. Epidermal growth factor receptor (EGFR, also known as erbB1 or HER-1), the member of the epidermal growth factor family (EGF-family), is vital for the proliferation and differentiation of OPCs [8-12]. After combined with its specific ligands, intrinsic intracellular protein-tyrosine kinase of EGFR auto-phosphorylated [13,14] and p-EGFR activated Akt1 [15,16], pAkt1 can not only up-regulate the expression of oligodendrocyte transcription factor (Olig2) [17], which plays critical and positive roles in the generation of OPCs [18-20], but also down-regulate the expression of caspase-3 that contributes to the apoptosis of OPCs [21-23]. Hence, we supposed that the protective effect of electro acupuncture on neural myelin sheaths after compressed spinal cord injury is mediated through activation of EGFR and its downstream proteins. However, this hypothesis should be certified by experiments.

Through observing before and after the treatment two groups of neurological function, the numbers and ultrastructure of axonal myelinated fibers, the expression of p-EGFR, pAkt1, Olig2 and caspase-3 circumference variations, we tried to elucidate the protective mechanism of EA on neural myelin sheaths in CSCI.

Materials and Methods

A total of 90 Sprague-Dawley (SD) rats weighing 250 g to 320 g were randomly divided into nine groups: normal groups (neither do compression spinal cord injury nor EA stimulation), EA groups (receiving electro acupuncture stimulation after decompression surgery) and control groups (without receiving electro acupuncture stimulation after decompression surgery) for 1 day after decompression, EA and control groups for 7 days after decompression, EA and control groups for 14 days after decompression, EA and control groups for 21 days after decompression (n=10/group). All rats were provided by the Experimental Animal Center of Chongqing Medical University. All experimental procedures were performed in accordance with the Animal Care and Ethics Committee and were approved by the Ministry of Science and Technology of the People’s Republic of China. The rats were housed in a 26°C room on a 12:12 dark/light cycle and were given enough food and water.

Surgical procedure

The model of compressed spinal cord injury (CSCI) was referred to the method designed by our team [24]. The rats were anesthetized with the intraperitoneal injection of 3.5% chloral hydrate and were subsequently fixed on a surgical table lying on back. The laminectomy was performed at L1 to expose the spinal cord, and the processed anterior and posterior arthrosis was excised without damaging the spinal cord. A custom-made device for chronic compressed spinal cord injury was immobilized the exposed dorsal surface of the spinal cord.

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After sutured, the nuts were exposed out of the skin. Motor, sensory, bowel and bladder functions were completely regular after woke up. From the first day after surgery, the screw was screwed into 1/4 turn (about 0.25 mm), and then twisted every 3-4 days until the double lower limb paralysis, incontinence (approximately 20 days). It is regarded as a successful model and the rats were then decompressed.

**Electro acupuncture**

For the EA group, electro acupuncture treatment was deduced from the first day of decompression until sacrifice. All EA was given at 9:00 a.m. to 11:00 a.m. every day. The rats were fastened for 30 min during EA administration without anesthetization [25] (Because of complete paralysis of hind limbs, the mouse can’t touch in the stimulation). In this study, we selected Zusanli (ST36) and Huantiao (GB30) acu-points to treat rats with CSCI based on two considerations. Firstly, these two acu-points are able to speed up the conduction of nerve impulses. Secondly, they are the two essential acu-points for the treatment of CSCI, prescribed by the Classics of the Traditional Chinese Medicine (TCM1). After cleaning the skin with alcohol, a pair of needle was connected with the output terminals of an EA apparatus (Model SDZ-II, Suzhou HuaTuo Medical Instrument Co., Ltd, Suzhou, China). Dense-sparse frequencies (60 Hz for 1.05 s and 2 Hz for 2.85 s, alternatively) were used for EA [26] and the intensity of the stimulation was applied for 30 min at 2 mA. Animals in control groups didn’t receive any treatment.

**Neurological function assessment**

Locomotor activity was examined based on Basso, Beattie, and Bresnahan (BBB) rating scale in an open field, according to indications of published articles by two independent observers in a double-blind manner [27-29]. A total of 21 scores from 0 (complete paralysis) to 21 (normal) were recorded after decompression of CSCI at 1st, 7th, 14th and 21st days.

**Luxol fast blue**

Mice were humanely euthanized and spinal cord were removed and fixed in 4% paraformaldehyde for 24 hours at 4°C. Sections from the center of the injury were selected along with sections from 0.5 cm to 0.7 cm rostral and caudal to the injury site. The tissues were then dehydrated successively in 10%, 20% and 30% sucrose solution, and embedded in optimal cutting temperature (OCT) compound. Ten micrometer thick (10 µm) transverse sections of the spinal cord were manufactured on a frozen slicer. Spinal cord sections were stained with Luxol fast blue at 60°C for 2 h. Sections were then rinsed in 95% ethanol. Differentiation was stopped by distilled water until unmyelinated tissue turned white. Twelve random micrographs from the lateral funiculus were obtained under an Olympus microscope (TEM; Hitachi-7500, Hitachi Ltd., Tokyo, Japan). These images were analyzed using the MIAS image analysis system to determine the ratio of myelin thickness to axon diameter (G-ratio) [30].

**Double-labeling immunofluorescence**

To identify p-EGFR, pAkt1, caspase-3, Olig2and NG2+ (the marker of oligodendrocyte precursor cells) co-expression in the spinal cord sections, we used the primary antibodies listed in Table 1. Tissue sections were rewarmed, rinsed and incubated in 5% donkey serum (Jackson Immuno Research, Lancaster, PA, USA) for 1 h at 37°C in a humidified atmosphere to permeabilise the tissue and block non-specific protein-protein interactions. The tissues were then incubated with the primary antibody overnight at +4°C. These tissue sections were rinsed again with 0.01mol/L PBS. The secondary antibody (red) was cy3 conjugated goat anti-rabbit IgG (H+L) used at a 1/200 dilution for 1.5h at 37°C in a humidified atmosphere in the dark. Alexafluor 488 goat anti-mouse IgG (H+L) used to label NG2 (green) at a 1/200 dilution for 1.5 h at 37°C in a humidified atmosphere in the dark. Nuclear dye (4',6-diamidino-2-phenylindole, 1:20; Bestbio Inc., China) was used to stain the cell nuclei (blue) for 5 min. The tissue sections were then washed and mounted in 50% glycerol dissolved in PBS. The samples were observed under a confocal microscope (Leica TCS SP2, Germany). All the digital images from lateral funiculus were captured in a double-blind manner from four random fields per section of the injured epicenter of the cross sections in the rats. The number of p-EGFR+-NG2, pAkt1+-NG2, Olig2+-NG2 and caspase-3+-NG2 per field was counted for analysis.

**Western blot**

Tissues soaked at 4°C in buffer containing 50 mmol/L ethylenediaminetetraacetic acid, 2 µg/mL of leupeptin, 2 µg/mL of pepstatin A, 2 mmol/L phenyl methyl sulfonyl fluoride, and 200 KIE/mL aprotinin were broken down mechanically using a blender. The homogenates were then centrifuged at 10,000×g for 20 min at 4°C. The supernatant was collected, and protein concentration was determined using a Bradford assay kit (Bio-Rad, Hercules, CA, USA). The proteins of the sample were separated using 10% SDS-PAGE and then transferred to poly vinylidenedifluoride membrane. Blotted membranes were incubated in 5% skim milk to block non-specific protein-protein interactions. For immunoblotting, the following primary antibodies were used: the polyclonal rabbit anti-olig2 antibody (1:1000; Abcam, Cambridge, UK, ab13847) for the monoclonal rabbit anti-EGFR antibody (1:1000; Abcam, Cambridge, UK, ab66138); the monoclonal rabbit anti-EGFR antibody (1:1000; Abcam, Cambridge, UK, ab52894); and the polyclonal rabbit anti-caspase3 antibody (1:1000; Abcam, Cambridge, UK, ab4976). The secondary antibodies were used alkaline phosphatase–conjugated anti-IgG antibodies (1:10000, Santa Cruz). Immuno reactive bands were visualized using microscopy (TEM; Hitachi-7500, Hitachi Ltd., Tokyo, Japan).

**Table 1:** Antibodies used in immune-fluorescence and western blot in this study.

<table>
<thead>
<tr>
<th>Antibody name</th>
<th>Manufacturer</th>
<th>Dilution</th>
<th>Host</th>
<th>Labelling</th>
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</thead>
<tbody>
<tr>
<td>Anti-EGFR antibody (ab15669)</td>
<td>Abcam, USA</td>
<td>1:200</td>
<td>rabbits</td>
<td>Epidermal growth factor receptor</td>
</tr>
<tr>
<td>Anti-pAkt1 antibody (ab6138)</td>
<td>Abcam, USA</td>
<td>1:200</td>
<td>rabbits</td>
<td>pAkt</td>
</tr>
<tr>
<td>Anti-olig2 antibody (ab81093)</td>
<td>Abcam, USA</td>
<td>1:200</td>
<td>rabbits</td>
<td>Oligodendrocyte transcription factor 2</td>
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<tr>
<td>Anti-active caspase3 antibody (ab13847)</td>
<td>Abcam, USA</td>
<td>1:200</td>
<td>rabbits</td>
<td>Active caspase-3</td>
</tr>
<tr>
<td>Anti-NG2 antibody (ab50009)</td>
<td>Abcam, USA</td>
<td>1:200</td>
<td>mouse</td>
<td>NG2</td>
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</tbody>
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a chemiluminescent substrate (Pierce Inc., Rockford, IL, USA). Western bands were quantified by gel densitometry (Bio-Rad). A ratio of protein–beta-actin for each sample was obtained (each point was repeated in triplicate).

Statistical Analysis

Statistical analyses were performed using GraphPad Prism 5.0 (GraphPad Software, Inc., USA). Differences between control groups and EA groups were compared using two independent samples t-test. Correlation between phosphor-EGFR and pAkt-1, Olig2, caspase-3 was then analyzed with pearson correlation. P<0.05 was considered statistically significant.

Results

Neurological function assessment

To evaluate the effect of EA on functional recovery, we assessed the locomotor skills in an open field using the Basso, Beattie, and Bresnahan (BBB) rating scale. Two independent samples t-test was performed: No statistically significant differences were found between control groups and EA groups at day 1 and day 7 (EA for 1 day group vs. control for 1 day group, p>0.05; EA for 7 days group vs. control for 7 days group, p>0.05; Figure 1); while the mean BBB scores in EA groups for 14 days and 21 days were remarkably increased when compared with its corresponding groups in control group (EA for 14 days group vs. control for 14 days group, p<0.01; EA for 21 days group vs. control for 21 days group, p<0.01, Figure 1). The results suggested that EA could protect the neurological function if we have enough time.

Demyelination after CSCI

Myelinated nerve fibers were investigated using luxol fast blue (LFB) and the number of myelinated nerve fibers was counted by Image J (National Institutes of Health, USA). Statistical analysis showed that no statistically significant differences were found between control groups and EA groups at 1st day and 7th day (EA for 1 day group vs. control for 1 day group, p>0.05; EA for 7 day group vs. control for 7 day group, p>0.05; Figure 2), while the mean LFB scores in EA groups for 14 days and 21 days were remarkably increased when compared with its corresponding groups in control group (EA for 14 days group vs. control for 14 days group, p<0.01; EA for 21 days group vs. control for 21 days group, p<0.05; Figure 2). These results suggested that EA have a protective effect on neural myelin sheaths.

Ultra-structural results

We then investigated the ultrastructural features of myelinated nerve fiber at the lateral funiculus in the white matter by transmission electron microscope (TEM). The result showed that axons had a varied swelling degree and cell organelles in the axoplasm became degenerative and lost both in control and EA groups, and the layers of myelin sheaths became disordered, thickened and even broken down after CSCI (Figure 2). However, the swelling degree of myelin sheaths in EA groups for 14th day was milder than those in control groups and the layers of myelin sheaths in EA groups for 14th day were more compact than those in the control groups (Figure 2).

Expression of p-EGFR and pAkt1

The p-EGFR+-NG2 and pAkt1+-NG2 cells in the spinal cord sections were identified by double-labeling immunofluorescence and the expression of p-EGFR and pAkt1 was detected by western blot, using antibodies against NG2, p-EGFR and pAkt1. Double-labeling immunofluorescence results showed that the p-EGFR+-NG2 and pAkt1+-NG2 cells was increased and reached the maximum on 14th day both in control and EA groups (Figures 3B-3E and Figures 4B-4E); the p-EGFR+-NG2 and pAkt1+-NG2 cells were distributed sparsely in control groups but widely in EA groups for 14th day (Figures 3C-3E and Figures 4C-4E); a positive correlation was also found between the number of p-EGFR+-NG2 and pAkt1+-NG2 cells (Not displayed in the picture). The western blot results showed that: compared with control group, the expression of p-EGFR and pAkt1 in EA groups for 7th, 14th and 21st day was significantly increased (Figures 3F-3H; EA for 14th day vs. control for 14th day, p<0.05; EA for 7th day vs. control for 7th day, p<0.05; EA for 21st day vs. control for 21st day, p<0.05). Figures 4F-4H, EA for 14th day vs. control for 14th day, p<0.05; EA for 7th day vs. control for 7th day, p<0.05; EA for 21st day vs. control for 21st day, p<0.05).

Expression of Olig2 involved in OPCs proliferation/differentiation

To further explore the protective mechanism of electroacupuncture, we examined Olig2 which is participated in oligodendrogenesis [31-34] and can be regulated by pAkt117. Double-labeling immunofluorescence results showed that: The Olig2+-NG2 cells was increased and reached the maximum on 14th day in control groups (Figures 5B-5E); the Olig2+-NG2 cells were distributed sparsely...
Figure 2: The pathological changes in myelinated nerve fibers at the lateral funiculus in the white matter were detected by luxol fast blue and transmission electron microscopy (TEM). (A-E) The myelinated nerve fibers at the lateral funiculus in the white matter in control groups, scale bar: 25 mm. (F-J) The myelinated nerve fibers at the lateral funiculus in the white matter in control groups, scale bar: 1 mm. (K-P) The ultrastructural features of myelinated nerve fibers at the lateral funiculus in the white matter in control and EA groups, scale bar: 1 mm. Extensively distributed myelinated fibers in normal groups were routinely observed and the myelinated axons showed a normal axoplasm with well-preserved cellular structure. The axons both in control and EA groups for 1st day varied swollen, myelin sheaths became edema, the layer of myelin sheaths became disorder, thick and even broken down. The swollen degree of myelin sheaths in control groups for 14th day was milder than that in control groups for 1st day but never return normal. The number of myelinated axons at the corresponding times both in control and EA groups. Compared with the normal groups, the number of myelinated nerve fibers was significantly decreased both in control and EA groups, and not returned to normal. No statistically significant differences were found between the control and EA groups for 1st day and 7th day (P>0.05). However, the number of myelinated nerve fibers in EA groups was remarkably increased compared with control groups for 14th and 21th day. (EA for 14 days group vs. control for 14 day group, P<0.05; EA for 21 days group vs. EA for 21 days group, P<0.05).

Figure 3: The number of p-EGFR+NG2 cells by double-labelling immunofluorescence and the expression of p-EGFR by Western Blot in different time point. A, B, C, D, E respectively represents the number and fluorescence intensity of p-EGFR+NG2 in normal, control for 1 day, control for 14 days, EA for 1 day and EA for 14 days’ group. F, G respectively represents the expression of p-EGFR in control and EA groups. H Data represent mean ± SEM (n=3). Two independent samples t-test showed that no statistically significant differences were found between control and EA groups at day 1; whereas, the expression of EGFR in EA groups for 7th, 14th and 21th day was significantly increased when compared with the corresponding group in control groups (P<0.05).
Figure 4: The number of pAkt1+–NG2 cells by double-labelling immunofluorescence and the expression of pAkt1 by Western Blot in different time point. A, B, C, D, E respectively represents the number and fluorescence intensity of pAkt1+–NG2 in normal, control for 1 day, control for 14 days, EA for 1 day and EA for 14 days’ group. F, G respectively represents the expression of pAkt1 in control and EA groups. H Data represent mean ± SEM (n=3). Two independent samples t-test showed that no statistically significant differences were found between control and EA groups at day 1: whereas, the expression of pAkt1 in EA groups for 7th, 14th and 21st day was significantly increased when compared with the corresponding group in control groups (P<0.05).

Figure 5: The number of Olig2+–NO2 cells by double-labelling immunofluorescence and the expression of Olig2 by Western Blot in different time point. A, B, C, D, E respectively represents the number and fluorescence intensity of Olig2+–NO2 in normal, control for 1 day, control for 14 days, EA for 1 day and EA for 14 days’ group. F, G respectively represents the expression of Olig2 in control and EA groups. H Data represent mean ± SEM (n=3). Two independent samples t-test showed that no statistically significant differences were found between control and EA groups at day 1; whereas, the expression of Olig2 in EA groups for 7th, 14th and 21st day was significantly increased when compared with the corresponding group in control groups (P<0.05).
in control groups but widely in EA groups for 14th day (Figures 5C-5E); a positive correlation was also found between the number of pAkt1+-NG2 and Olig2+-NG2 cells (Not displayed in the picture). Western blot results showed that: the expression of Olig2 in EA groups for 7th, 14th and 21th day was significantly decreased when compared with its corresponding group in control groups (p<0.06).

Expression of caspase-3 involved in apoptosis

Caspase-3 has been shown to play central role in apoptosis pathways. Meanwhile, the expression of caspase-3 can be down-regulated by pAkt1. Therefore, we examined the expression of caspase-3. Double-labeling immunofluorescence results showed that: the caspase-3+NG2 cells was abruptly increased at day 1 and then decreased with time both in control and EA groups (Figures 6B-6E); caspase-3+NG2 cells were distributed widely in control groups but sparsely in EA groups for 14th day (Figures 6C-6E); a negative correlation was also found between the number of pAkt1+NG2 and caspase-3+NG2 cells (Not displayed in the picture). The western blot results showed that: the expression of caspase-3 was abruptly increased at day 1 (control for 1st day vs. normal, p<0.01; EA for 1st day vs. normal, p<0.01) and then decreased with time both in control and EA groups. Furthermore, the expression of caspase-3 in EA group for 7th, 14th and 21th day was significantly decreased when compared with its corresponding group in control groups (p<0.05).

Figure 6: The number of caspase-3+NG2 cells by double-labelling immunofluorescence and the expression of caspase-3 by Western Blot in different time point. A, B, C, D, E respectively represents the number and fluorescence intensity of caspase-3+NG2 in normal, control for 1 day, control for 14 days, EA for 1 day and EA for 14 days’ group. F, G respectively represents the expression of caspase-3 in control and EA groups. H Data represent mean ± SEM (n=3). The expression of caspase-3 was abruptly increased at day 1 (control for 1 st day vs. normal, p<0.01; EA for 1 st day vs. normal, p<0.01) and then decreased with time both in control and EA groups.

Discussion

Previous studies reported that EA treatment can increase the number and differentiation of endogenous OPCs and remyelination in the demyelinated spinal cord. However, the specific mechanism of promoting the proliferation and differentiation of endogenous OPCs and remyelination is largely unknown. In addition, it is known that extensive demyelination induced by oligodendrocytes apoptosis was the major pathological change after CSCI. Therefore, the independent designing CSCI model was developed to observe whether electroacupuncture could promote the proliferation and differentiation of OPCs, remyelination and functional recovery. After 7- day treatment of EA, the neurological function was significantly improved, consistent with the improved ultrastructural features and numbers of myelinated nerve fibers. This result is consistent with the previous studies, which further shows that electro acupuncture can promote the proliferation and differentiation of OPCs, myelin formation and neural function recovery.

In order to further explore the protective mechanism of electroacupuncture on neural myelin sheaths, we next investigated the expression of p-EGFR and its downstream protein—pAkt1. Significant studies have been reported that p-EGFR functions as a regulator in the oligodendrocyte lineage, including induction of survival and proliferation. After activated by specific ligands, such as EGF or neuregulin, EGFR phosphorylated and bound with SH2 domains of pAkt1, thereby regulating the expression of other proteins in the cell. In this study, we found that the expression of p-EGFR, pAkt1 in the EA groups was significantly increased and p-EGFR-NG2+, pAkt1-NG2+ were found widely in the white matter. These results are consistent with the changes of neuroglial function, ultrastructural features and numbers of myelinated nerve fibers. The result suggested

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that p-EGFR, pAkt1 were clearly up-regulated by EA to improve the recovery of the injured spinal cord.

Akt1 is one of the members of the p-Akt (Protein kinase B, PKB) family, which is a key signaling protein in the cellular pathways. It can not only can not only up-regulate the expression of oligodendrocyte transcription factor (Olig2) [17], but also down-regulate caspases family [39,40]. Therefore, we have tested Olig2 and caspase-3 based on the test of pAkt1.

The formation of oligodendrocytes is triggered by an increased activity of bHLH factors, in which Olig2 coupled with an increase in pAkt1 activity [17]. Olig2 is a class of bHLH transcription factor that is specially expressed by Olig cells in a restricted domain of the spinal cord ventricular zone and sequentially generates oligodendrocytes [41]. Olig2 is excellent in positively controlling OPCs generation [42-44] and can strongly up-regulate the OPCs in demyelinating diseases to benefit neuronal repair [45]. In this study, we found that the expression of Olig2 in the EA groups was significantly increased and Olig2+-NG2 cells were widely distributed in the white matter. Meanwhile, the increase of Olig2 is consistent with the up-regulation of pAkt1. The result suggested that Olig2 was clearly up-regulated by EA to promote the recovery of the injured spinal cord and this effect may be attributed to pAkt1.

Caspases are a family of protease enzymes playing essential roles in programmed cell death, in which Caspase-3 that mediated the recovery of the injured spinal cord and this effect may be attributed to pAkt1. The main apoptosis pathways, including death receptor pathway, mitochondrial pathway and endoplasmic reticulum stress (ER stress) pathway [46], has been shown to play a central role. Our result also revealed that the expression of caspase-3 was even increased after CSCI, suggesting that caspase-3 plays a vital role in apoptosis of oligodendrocytes after CSCI. Meanwhile, the expression of caspase-3 can be down-regulated by pAkt [47]. Hence, we next investigated the expression of caspase-3. In the current study, we found that the expression of caspase-3 in the EA groups was significantly decreased and caspase-3+-NG2 cells were sparsely distributed in the white matter. And the expression of caspase-3 and pAkt1 was negatively correlated: when the expression of pAkt1 was increased, the expression of Caspase-3 was decreased, and vice versa. These results suggested that EA stimulation can prevent oligodendrocytes from apoptosis by down-regulating the expression of caspase-3 and this effect may be closely related to the up-regulation of pAkt1.

Conclusion

This study showed that the protective effect of EA on neural myelin sheaths might be mediated via activation of EGFR and its downstream associated proteins after CSCI. This study can provide experimental basis for the clinical application of EA.

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Declaration of Rights

The authors state that there are no actual or potential conflicts of interests.

Submission Declaration and Verification

The authors have not published or submitted the manuscript elsewhere.

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