Research Article
Effects of Electroacupuncture at Governor Vessel Acupoints on Neurotrophin-3 in Rats with Experimental Spinal Cord Injury

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1. Introduction

Spinal cord injury (SCI) is an accidental tragedy, causing unexpected suffering physically and emotionally, and is costly to patients [1]. Traumatic SCI can cause disorders of somesthesia and locomotion below the level of injury. SCI induces primary mechanical damage and causes secondary damage to the spinal cord. Primary damage occurs by mechanical tissue disruption immediately subsequent to trauma. Secondary damage is mediated by complex cellular and molecular processes [2].

At present, mainstream treatments for SCI include pharmacotherapies, neurotrophic factors, cell-transplantation, gene therapy, and biological materials transplantation [3, 4]. Traditional Chinese Medicine (TCM) treatments were also reported and used [3, 4]. Pharmacotherapy includes...
methylprednisolone and GM1 [3]. Neurotrophic factors include nerve growth factor (NGF) and brain-derived neurotrophic factor (BDNF) [5].

Acupuncture is a therapeutic technique used in TCM. Since its development several thousand years ago, acupuncture has made many contributions to health care and medical treatment. Electroacupuncture (EA) is a type of therapy in which a needle inserted into an acupoint is attached to a trace pulse current with the purpose of producing a synthetic electric needling stimulation. The application of EA for the treatment of SCI has shown promising results in the alleviation of the patients' suffering [6]. Previous studies have shown that applications of EA for the treatment of SCI have been proven to contribute to neurologic and functional recoveries in SCI [7–9].

TCM believes that Governor Vessel injury is the main reason of SCI. Malnutrition of tendons and muscles due to obstruction of qi and blood induces disuse and atrophy. Therefore, Governor Vessel acupoints are the first choices for treating SCI. “Dazhui” (GV 14) is the confluence of Governor Vessel and the hand and foot three Yang meridians. Acupuncture stimulation in GV 14 can activate and inspire Yang-qi in the whole body to dredge the meridians. “Mingmen” (GV 4) is also a Governor Vessel acupoint and is the intersection of the Governor Vessel and Belt Vessel. It gathers genuine-Yin and genuine-Yang, which are the root of Yuan-qi and the gateway of life. Stimulating GV 4 can regulate channels and activate collaterals, tonify Yang, and strengthen the kidneys. In previous experiments using SCI rat models, acupuncture stimulation in GV 14 and GV 4 has been adopted [10].

EA on the Governor Vessel has been shown to alleviate secondary damage after SCI in both patients and animal models [10, 11]. Therapeutic effects has been reported by a review [12] which showed that acupuncture stimulation on Governor Vessel could promote motor function recovery after SCI and improve restoration of bladder function. Evidence from both clinical trials and basic researches supports EA on Governor Vessel for restoration of motor function, bladder function, and sensory function of limbs after SCI [10, 13–15].

Neurotrophic factors (NTFs) are protein molecules which are essential to neuron survival and growth [16]. NGF, BDNF, neurotrophin 3 (NT-3), and neurotrophin 4/5 (NT-4/5) have been determined as NTFs. NT-3 is crucial to neuron survival, differentiation, and formation of neural circuits during neural development [17, 18]. NT-3 is also an important factor in the microenvironment for spinal cord repair [16]. It plays an influential role in preventing neuron death in injured spinal cord, maintaining neuron survival and promoting axon regeneration [16]. Exogenous NT-3 also promotes adult stem cell survival and differentiation after transplantation into an injured spinal cord [19]. Previous research found that EA on Governor Vessel promoted secretion of NT-3 at the injured area and adjacent tissues 14 days after SCI [7, 20]. However, whether NT-3 is altered prior to this time is not known.

This study aims to explore whether EA can promote a suitable microenvironment for recovery of neurological function after SCI by increasing endogenous NT-3 secretion and expression at the injured area and adjacent tissues after performing EA on the Governor Vessel during the early phase of SCI.

2. Experimental Procedures

2.1. Animal and Experimental Groups. All experiments were approved by the Institutional Animal Care and Use Committee of Beijing University of Chinese Medicine. 165 rats (Sprague-Dawley, male, 180 to 220 g) were randomly assigned into three groups at a ratio of 1:1:1. The sham-operation group (S) received only a laminectomy. The remaining two groups were modeled as spinal cord injury at the T10 spinal segment. The control group (C) received no treatment. The EA group received EA treatment at GV 14 and GV 4 acupoints. Rats in the control group and EA group were randomly placed into the following subgroups: 1d, 3d, and 7d. There were 20 rats per group in 3 d and 7 d, respectively, and 15 rats in group 1d. Each of the animals was housed in separated cages with free access to food and water. Room temperature was set at 25 ± 3°C.

2.2. Spinal Cord Injury. The surgical procedure for inducing SCI was conducted according to established methods [21]. The rats were anesthetized with 10% chloral hydrate (3.5 mg/kg, intraperitoneal), and a laminectomy was performed at the T9–T11 level, exposing the underlying cord without disrupting the dura. Spinal cord contusion was induced using a weight-drop apparatus, where a guided 5 g rod was dropped from a height of 80 mm onto the exposed cord, representing moderate SCI. The absorbable gelatin sponge was placed at the site of the SCI for hemostasis and was sutured to the vertebral column. Afterwards, the skin and musculature were sutured. Laminectomy was performed on the sham-operation group without SCI. All SCI animal models included in the study met the following injury criteria: spinal cord ischemia and edema around the wound, formation of tail sway reflex, flicking of both body and legs, and the appearance of sluggish paralysis (Figure 1). All the animals were returned to separated cages with sufficient water and food and were then treated everyday with an intraperitoneal injection of gentamicin at a dose of 2000 μ, twice daily. No analgesics were used. Rats that underwent SCI received specialized care consisting of manual bladder expression three times daily and cleansing for the duration of the experiment.

2.3. Acupuncture Application. GV 14 and GV 4 acupoints were utilized during EA treatment. The GV 14 acupoint is located in the posterior midline and in the depression below the spinous process of the 7th cervical vertebra in prone position. GV 4 is also located in the posterior midline and in the depression below the spinous process of the 2nd lumbar vertebra in prone position. Rats were placed within a cloth bag without anesthesia during EA administration. Sham and control animals were captured for bondage when the EA group received treatment, for a duration of 20 minutes each time, to ensure the same processing conditions. Sterilized
disposable stainless steel needles (0.30 × 25 mm, Zhongyan Taihe brand, Beijing Zhongyan Taihe Medical Instrument Co., Ltd., Beijing, China) were inserted into GV 14 (oblique downwardly) and GV 4 (oblique upwardly), respectively, as deep as 5–7 mm for both points. Following the insertions, electrodes were connected to the handle of the needles (electric acupuncture apparatus used: LH202 acupoint nerve stimulator, Beijing Huawei Industrial Development Co., Ltd., Beijing, China). Electric simulation parameters were at a frequency of 2 Hz and an intensity of 1 mA for 20 minutes. The rats received EA treatment 2 hours after the SCI model was established and anesthesia recovery. One hour before euthanasia, at the 24th hour of postsurgery, the rats in 1 d EA group were given the 2nd treatment. The rats in 3 d and 7 d EA group were given the same treatment once per day until they were euthanized at the appointed time. The sham and control animals did not receive treatment. Finally, 11 rats per group were analyzed. 6 rats per group were perfused to fixation and fresh tissue was taken from the other 5 rats per group. All the above procedures were conducted after anesthesia.

2.4. Behavioral Testing. Rats from the sham group, the control group, and the EA group were assessed for hind limb motor function at 1 d, 3 d, and 7 d after injury by two blinded observers using the Basso, Beattie, and Bresnahan (BBB) hind limb locomotor rating scale test. The BBB rating scale is a 21-point system based on operationally defined behavioral features, which follow the recovery progression from complete paralysis to normal locomotion. The rating scale ranges from 0 to 21, with a score of 0 indicating complete hind limb paralysis and a score of 21 denoting completely normal locomotor function.

2.5. Nissl Staining. Six of eleven rats in each group were euthanized at the specified time of postsurgery and perfused transcardially with 0.9% sodium chloride and then with 4% paraformaldehyde in 1× phosphate buffered saline (4% PFA) for 30 min. Spinal cords were surgically removed and kept in 4% PFA for postfix overnight. After dehydration, the spinal cords were embedded with paraffin, and serial coronal sections with thickness of 4 µm were obtained. To assess the histopathologic change, each one of six sections obtained was subjected to Nissl staining. Surviving neurons were characterized by blue staining Nissl bodies. Quantification was performed by counting the quantity of survival neurons in five randomly chosen fields within each slide from each section in the same part of the structure at 400x with an Olympus (BX53, Japan) optic microscope. Two observers, blind to the experiment, counted the surviving neurons.

2.6. TUNEL Staining. To detect apoptosis, each one of six coronal sections obtained in the above experimental procedure was further subjected to terminal deoxynucleotidyl
transferase mediated dUTP nick end labeling (TUNEL) staining. To assess the apoptotic cells, we measured 6 samples in one section chosen at random from each spinal cord. Apoptotic cells were characterized by dark brown staining of nucleus and nuclear membrane. Quantification was performed by counting the quantity of positive cells in five randomly chosen fields within each slide from each section in the same part of the structure at 400x with an Olympus (BX53, Japan) optic microscope. The index of apoptosis was calculated as the ratio of overall apoptotic cells. Two observers, blind to the experiment, counted the apoptotic cells.

2.7. Western Blotting. The remaining five of eleven rats in each group were deeply anesthetized with 10% chloral hydrate and euthanized at the appointed time. A 1 cm spinal cord segment centered at the injury epicenter was quickly dissected. Spinal cord protein homogenates were prepared by rapid homogenization in a 50% sucrose solution at 4°C. Protein concentrations were determined using the Bradford method [22]. For electrophoresis, protein samples (40 μg each) were dissolved in the sample buffer and heated to 100°C for 5 min. Samples were then resolved on 10% SDS-PAGE and transferred to PVDF membranes. The membranes were blocked in TBS-T containing 5% nonfat dry milk for 1 h, followed by incubation with NT-3 (1:500, ABcam, UK) at 4°C overnight. After being washed, the membranes were incubated with HRP-Goat-Anti-Mouse IgG (1:5000, ZSGB-BIO, Beijing, China). Proteins were visualized by ECL Chemiluminescence (Santa Cruz, USA). All Western blot experiments were repeated for at least 3 times. GAPDH (reduced glyceraldehyde-phosphate dehydrogenase) served as an internal control. The gray values of target proteins were divided by that of internal control to correct the error, which resulted due to the relative content of the target protein in the sample.

2.8. Statistical Analyses. No rats died in 1d; while in 3d, 2 rats died in the sham group, 3 rats died in the control group, and 1 rat died in the EA group; in 7d, no rats died in the sham group, 7 rats died in the control group, and 5 rats died in the EA group. Lastly, 11 rats per group were randomly selected to enter into statistics. Data were presented as means ± SD (Standard Deviation). SPSS 20.0 (SPSS Inc., Chicago, USA) was deployed for data analysis with a one-way ANOVA method after the test of normal distribution and homogeneity of variance, followed by a post hoc multiple comparison. Between the two groups, we used Fisher’s LSD (Least Significant Difference) method to compare any differences. Statistical significance was set to $p < 0.05$.

3. Results

3.1. Effects of EA on Behavioral Testing after SCI. We used the BBB hind limb locomotor rating scale test to assess neurological function at 1d, 3d, and 7d after SCI. The mean BBB scores (Table 1) of the control group were lower than the sham group at 1d, 3d, and 7d after SCI ($p < 0.01$). The mean BBB scores in the control group increased from 0.6364 at 1d to 2.0909 at 7d (Table 1). The mean BBB scores in the EA group increased from 0.6364 at 1d to 4.0909 at 7d (Table 1). The BBB scores of the sham group were at the highest level because nerve function was not damaged [23]. These results suggested that EA plays an important role in improvement of neurological function.

2.8. Statistical Analyses. No rats died in 1d; while in 3d, 2 rats died in the sham group, 3 rats died in the control group, and 1 rat died in the EA group; in 7d, no rats died in the sham group, 7 rats died in the control group, and 5 rats died in the EA group. Lastly, 11 rats per group were randomly selected to enter into statistics. Data were presented as means ± SD (Standard Deviation). SPSS 20.0 (SPSS Inc., Chicago, USA) was deployed for data analysis with a one-way ANOVA method after the test of normal distribution and homogeneity of variance, followed by a post hoc multiple comparison. Between the two groups, we used Fisher’s LSD (Least Significant Difference) method to compare any differences. Statistical significance was set to $p < 0.05$.

3.2. Effects of EA on Neuron Survival after SCI. In the sham group, neurons exhibited a large amount of densely stained toluidine blue granules in the cytoplasm (Figure 2) [24]. However, in the control group, the Nissl bodies dramatically decreased or even disappeared in the neurons at 1d (Figure 2). At 3d, the motor neurons of the gray matter showed obvious atrophy, some neurons dissolved, and the quantity of the Nissl bodies was significantly reduced or even disappeared in the control group (Figure 2). At 7d, a small number of neurons’ Nissl bodies appeared again in the control group (Figure 2). In the EA group the quantity of Nissl bodies was restored compared with that of the control group and displayed patch morphology [24]. The histomorphology had no change in EA group at 1d; neurons also had varying degrees of reducing as dyeing was lighter (Figure 2). At 3d, neuronal atrophy and reduction still existed, but the condition was better than in the control group, and Nissl bodies were faintly visible in the EA group (Figure 2). At 7d, the Nissl bodies of the remaining neurons appeared again after receiving EA treatment, and the quantity was significantly larger than that of the control group (Figure 2). EA group showed significantly preserved neurons compared with that of the control group at 1d, 3d, and 7d after SCI (Figure 2; Table 2) (data are presented as mean ± SD). **$p < 0.01$, versus sham group; *$p < 0.05$, versus control group; $n = 6$ animals per group). These results suggested that rats had a lower number of surviving neurons after SCI, and EA can improve survival.

3.3. Effects of EA on the Inhibition of Apoptotic Cell Death after SCI. In the sham group, there were almost no TUNEL positive cells (Figure 3). In the control group, positive cells appeared and mainly distributed in the damaged area and its edge at 1d after SCI (Figure 3). The positive cells significantly increased and widely distributed in both the white and gray matter of the control group at 3d after SCI (Figure 3).

| Group | 1d | 3d | 7d |
|-------|----|----|----|
| S     | 21 ± 0.00 | 21 ± 0.00 | 21 ± 0.00 |
| C     | 0.6364 ± 0.15** | 1.3636 ± 0.15** | 2.0909 ± 0.21** |
| EA    | 0.6364 ± 0.15** | 2.0909 ± 0.25** | 4.0909 ± 0.21**## |

*Table 1: Behavioral testing after SCI in the following groups ($n = 11$ per group): sham, control, and EA. **$p < 0.01$, as compared with sham group; *$p < 0.01$, as compared with control group (S: sham group; C: control group; EA: electroacupuncture group).*
Neural Plasticity

Figure 2: Neuronal conditions were evaluated using Nissl staining (original magnification ×400, 1: 50) in the following groups: sham, control, and EA group at 1 d, 3 d, and 7 d (S: sham group; C: control group; EA: electroacupuncture group).

Table 2: Number of surviving neurons after SCI in the following groups (n = 6 per group): sham, control, and EA. ** p < 0.01, as compared with sham group; * p < 0.05; ** p < 0.01, as compared with control group (S: sham group; C: control group; EA: electroacupuncture group).

| Group | Number of survival neurons |
|-------|---------------------------|
|       | 1d                        | 3d                        | 7d                        |
| S     | 39.67 ± 1.54              | 38.50 ± 1.34              | 37.33 ± 1.15              |
| C     | 9.33 ± 1.12**             | 16.50 ± 1.12**            | 19.00 ± 0.97**            |
| EA    | 19.33 ± 1.52***           | 21.00 ± 1.07***           | 27.50 ± 1.38***           |

[24]. At 7 d, the positive cells were less than before in the control group. In the control group, neurons shrank and exhibited abnormal morphology with condensed chromatin, introverted nuclear membranes, and increased apoptotic bodies [24]. Neurons in the EA group, however, showed less condensed chromatin and clear nuclear membranes [24]. EA group, however, showed significantly decreased brown-positive cells compared with that of the control group at 1 d, 3 d, and 7 d after SCI (Figure 3; Table 3) (data are presented as mean ± SD, ** p < 0.01, versus sham group; * * p < 0.01, versus control group; n = 6 animals per group). This suggested that EA can reduce apoptosis, promote the survival of neurons, and has a certain protective effect.

3.4. EA Regulate Protein Expression of NT-3 after SCI. We used Western blotting analyses to determine the expression of NT-3 semiquantitatively for assessing the progression of SCI and the protective effects of EA. In the sham group, NT-3 demonstrated expression. In the control group, the expression of NT-3 decreased at 1 d and 3 d without statistical significance compared with the sham group (Figure 3; Table 3) (data are presented as mean ± SD, ** p < 0.01, versus sham group; * * p < 0.01, as compared with sham group; * * * p < 0.01, versus control group; n = 6 animals per group). This suggested that EA can reduce apoptosis, promote the survival of neurons, and has a certain protective effect.

Table 3: The effects of EA on the inhibition of apoptotic cell death after SCI in the following groups (n = 6 per group): sham, control, and EA. ** p < 0.01, as compared with sham group; * * * p < 0.01, as compared with control group (S: sham group; C: control group; EA: electroacupuncture group).

| Group | Apoptotic index |
|-------|----------------|
|       | 1d            | 3d            | 7d            |
| S     | 4.00 ± 1.87   | 5.00 ± 1.73   | 4.30 ± 1.15   |
| C     | 27.40 ± 3.9** | 29.00 ± 4.58**| 20.00 ± 1.00**|
| EA    | 14.00 ± 1.58***| 14.00 ± 1.00***| 12.60 ± 2.08***|
Figure 3: TUNEL staining (original magnification ×400, 1:50) identified apoptotic neurons in the following groups: sham, control, and EA group at 1 d, 3 d, and 7 d (S: sham group; C: control group; EA: electroacupuncture group).

Table 4: The effects of EA on the expression of NT-3 after SCI in the following groups (n = 5 per group): sham, control, and EA. *p < 0.05, as compared with sham group; **p < 0.01, as compared with control group (S: sham group; C: control group; EA: electroacupuncture group).

| Group | Relative expression of NT-3 |
|-------|-----------------------------|
|       | 1 d            | 3 d            | 7 d            |
| S     | 0.88 ± 0.21    | 1.09 ± 0.27    | 1.00 ± 0.08    |
| C     | 0.67 ± 0.14    | 0.60 ± 0.02    | 0.64 ± 0.28*   |
| EA    | 0.80 ± 0.21    | 0.98 ± 0.05**  | 1.10 ± 0.25**  |

4. Discussion

In this research, the BBB scores of control group and EA group which received SCI operation had statistical difference compared with that of sham group in 1 d, 3 d, and 7 d, which proved that the models of SCI were successfully established as the motor function of rats’ hind limbs appeared paralyzed. At 7 d after SCI, the BBB scores began to show significant statistical difference between EA group and the control group, which indicated that the hind limbs’ motor function of rats was gradually recovering, and the recovery of rats that received EA treatment was better than that of control rats without EA treatment. This signifies that EA treatment could promote motor function recovery after SCI, which was also shown by Nissl staining, TUNEL, and Western blotting.

In addition to the damage at the site of the spinal cord injury, secondary pathological changes occur in the following order: edema, ischemia, calcium overload, lipid
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NT-3 → 1 2 3 → 29 kDa
GAPDH → 1 2 3 → 36 kDa
NT-3 → 1 2 3 → 29 kDa
GAPDH → 1 2 3 → 36 kDa
NT-3 → 1 2 3 → 29 kDa
GAPDH → 1 2 3 → 36 kDa

Figure 4: Western blotting identified the expression of NT-3 in the following groups: sham, control, and EA group at 1 d, 3 d, and 7 d (1: sham group; 2: control group; 3: electroacupuncture group).

peroxidation, microcirculation obstruction, and apoptosis [25, 26]. Unlike neurons of the peripheral nervous system (PNS), those of the central nervous system (CNS) do not spontaneously regenerate their axons following injury; hence, trauma to the brain or spinal cord frequently results in permanent neurological deficits [27]. SCI induced apoptotic cell death of neurons and oligodendrocytes has been known to cause progressive degeneration of the spinal cord, leading to permanent functional deficits [28]. The development of effective therapies that can restore lost neurological function is therefore critically dependent on strategies to promote robust axon regeneration [27]. Over the past several decades, a great body of work has chipped away at elucidating the biological mechanisms limiting the regeneration of injured CNS axons [27].

Our results showed that less postinjury neuronal death was occurring in the EA group compared with the control group, which was interpreted as neuroprotection. Furthermore, our results still indicated that EA could maintain the cell morphology of neurons, which contributed to the functional recovery of the injured spinal cords of the rats.

Neurotrophic factors enter axons by way of its receptor mediated endocytosis at the end of axons, then reach neurons through plasma flow reverse transportation, which can cause different signaling pathway activation or inhibition in the cell, and regulate relative protein expression, to play its effect of supporting neuron development, survival, growth, and function integrity [16]. NT-3 is crucial to neuron survival, differentiation, and formation of neural circuits during neural development [17, 18].

NT-3 is widely distributed in peripheral and central nervous systems, mainly in the hippocampus, dorsal root ganglia, brainstem, and spinal cord et cetera [29]. Chen and Fang found that NT-3 like immunoreactive substance in the distribution of central nervous system involves both glial cells and neurons; the former is mainly distributed in the corpus callosum, substantia nigra, fimbria of hippocampus, subependymal zone, and cerebellum, while the latter is mainly distributed in the septum, diagonal band of Braco, granulosa cells of primitive olfactory cortex, amygdala, brainstem, and spinal motor neurons [29]. This could be understood as the gray matter of the spinal cord as the main expression site of NT-3, particularly expressed in motor neurons, so NT-3 for neurons in the gray matter of the spinal cord, especially in ventral horn motor neurons’ survival and normal function, played an important role [29–31].

The present study found that the level of NT-3 in the injured spinal cords of the control group and EA group decreased in comparison with that in sham group rats’ spinal cord where it was still synthesized and secreted. This might be injured spinal cord induced cell death or the decline of damaged cells’ function in the damaged region and adjacent tissues which synthesize and secrete NT-3; therefore the NT-3 levels would be decreased [32]. We also found the level of NT-3 in the spinal cord injury region and adjacent tissues of EA group significantly increased compared to that of control group after receiving 7 days of Governor Vessel EA treatment. This might be induced by Governor Vessel EA treatment preventing the death of some damaged cells which synthesize and secrete NT-3 or possibly the regulation and restoration of the damaged cells’ physiological function. No matter what the mechanism of increased NT-3 was, the increased level of NT-3 in the region of spinal cord injured and adjacent tissues created an appropriate microenvironment for spinal cord repair. It was shown in the following research results that Chen et al. [33] used immunohistochemical staining, in situ hybridization, and PCR technique to detect and observe the influence of EA for the expression of NT-3 in the spinal cord dorsal root. The study found that the expression of NT-3 in large or small neurons in cats’ dorsal root ganglion of EA group was significantly higher than the other groups. Wang et al. [34] detected the treatment effect of EA through using a model of cats’ spinal cord dorsal root partial resection and found that the positive neurons of NT-3 in the spinal cord lamina II increased and speculated that EA could accelerate the repair of a spinal cord injury through promoting the expression of NT-3.

In addition, NT-3 plays important roles in oligodendrocyte development [35, 36]. Huang et al.’s study indicated that EA treatment could promote NT-3 expression and increase the number and differentiation of endogenous oligodendrocyte precursor cells (OPCs) and remyelination in the demyelinated spinal cord [7]. Another study showed that EA treatment could increase NT-3 expression and promote oligodendrocyte-like cell differentiation from (NT-3) receptor (TrkC) gene modified mesenchymal stem cells (TrkC-MSCs), remyelination, and functional improvement of demyelinated spinal cord. However in our study, we did not observe EA effects on NT-3 and its action on oligodendrocytes and other glial cells [37].
According to TCM, GV 14 and GV 4 are points pertaining to the Governor Vessel. The spinal cord and Governor Vessel have direct contact on channels and collaterals. Based on the principle of “where meridians pass, indications for acupoints of this meridian could be considered,” selecting the Governor Vessel to treat paraplegia related to spinal cord injury conforms to the TCM saying “searching for the primary cause of disease in treatment” [38, 39].

5. Conclusion

Governor Vessel EA on GV 14 and GV 4 could improve functional recovery by reducing apoptotic cell death after SCI. Meanwhile, the neuroprotective effects of EA treatment might be mediated by the level of NT-3 in the microenvironment around the spinal cord after injury. In addition, the present study suggested that stimulating GV 14 and GV 4, especially with electroacupuncture, might be an effective therapeutic strategy in spinal cord injury.

Additional Points

Limitations. This study has a small sample size. And the interaction among NT-3 and its receptor (such as TrkC) and the effect of EA on them at different times should be explored in the future. Furthermore, the EA-N1-3 effects on oligodendrocytes and other glial cells should be considered rather than neurons only. Lastly, quantitative analysis is also necessary, such as ELISA.

Competing Interests

The authors declare that there are no competing interests.

Authors’ Contributions

Yu-ping Mo and Hai-jiang Yao contributed equally to this paper. This paper was written by Yu-ping Mo and revised by Hai-jiang Yao. Yu-ping Mo performed EA treatment and wrote the completed paper. Hai-jiang Yao, Wei Lv, and Hong-tao Song were in charge of model establishment and sample collection. The Nissl staining, TUNEL, and Western blotting were performed by Ying-qi Mao and Xiao-chen Yuan, respectively. Liang-yu Song and Quan-kai Jing completed behavioral assessment and data extraction. Su-hua Shi performed data analysis and Zhi-gang Li arbitrated any disagreement and ensured that no errors occurred during the study. All authors contributed to the conception of the study and approved the publication of this paper.

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