Application of the chronic constriction injury of the partial sciatic nerve model to assess acupuncture analgesia

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Purpose: To validate and explore the application of a rat model of chronic constriction injury to the partial sciatic nerve in investigation of acupuncture analgesia.

Methods: Chronic constriction injury of the sciatic nerve (CCI) and chronic constriction injury of the partial sciatic nerve (CCIp) models were generated by ligating either the sciatic nerve trunk or its branches in rats. Both models were evaluated via paw mechanical withdrawal latency (PMWL), paw mechanical withdrawal threshold (PMWT), nociceptive reflex-induced electromyogram (C-fiber reflex EMG), and dorsal root ganglion immunohistochemistry. Electroacupuncture (EA) was performed at GB30 to study the analgesic effects on neuropathic pain and the underlying mechanisms.

Results: Following ligation of the common peroneal and tibial nerves, CCIp rats exhibited hindlimb dysfunction, hind paw shrinkage and lameness, mirroring those of CCI rats (generated by ligating the sciatic nerve trunk). Compared to presurgery measurements, CCIp and CCI modeling significantly decreased the PMWL and PMWT. EA at GB30 increased the PMWL and PMWT in both CCI and CCIp rats. Calcitonin gene-related polypeptide and substance P expressions were apparently increased in both CCI and CCIp groups, but were not different from each other. The C-fiber reflex EMG of the biceps femoris was preserved in CCIp rats, but it could not be recorded in CCI rats on the 5th day after nerve ligation. The C-fiber reflex EMG was reduced at 0, 1, and 2 minutes after EA in CCIp rats, but only at 0 and 1 minute after EA in normal rats.

Conclusion: The CCIp model is better than the CCI model for studying acupuncture analgesia on chronic neuropathic pain and the underlying mechanisms.

Keywords: neuropathic pain, chronic constriction injury, acupuncture, analgesia, C-fiber reflex

Introduction

Neuropathic pain, defined by the International Association for the Study of Pain as “pain caused by a lesion or disease of the somatosensory system”,1 is a pain syndrome characterized by spontaneous pain, hyperalgesia, allodynia, and sensory abnormalities.2 Sciatica, one of the most common sources of clinical neuropathic pain, is generally defined as pain radiating to the leg, normally below the knee and into the foot and toes.3 The treatment for sciatica remains a current challenge in clinic. Therefore, generating appropriate animal models are critical to developing novel treatment options for neuropathic pain.1–6

Chronic constriction injury of the sciatic nerve (CCI) was established in 1988 as a means to produce neuropathic pain in laboratory animal models;7 the pathological condition of this model is quite similar to human peripheral nerve injury and its consequent...
behavioral manifestations. Based on the CCI model, a variety of animal models have since been developed for different research purposes, including the partial sciatic nerve ligation (pSNL), the spinal nerve ligation to sciatic nerve (SNL), the sciatic nerve axotomy, the chronic compression of the dorsal root ganglion, and the spared nerve injury (SNI) models.

Although basic and clinical studies have confirmed that acupuncture can effectively alleviate or cure pain syndromes, the mechanisms responsible for acupuncture analgesia remain elusive. This suggests the need for a proper animal model in order to elucidate mechanisms by which acupuncture alleviates neuropathic pain. Behavior test and nociceptive stimulation-induced EMG (C-fiber reflex EMG) are both measures for pain alleviating evaluation. Paw mechanical withdrawal latency and threshold (PMWL and PMWT, respectively) are the tests of mechanical nociception or sensibility based on the rat’s sensory–motor reflex. It is withdrawal of the paw with participation of the upper central nervous system and multisynaptic relay. Whereas C-fiber reflex is a neuromuscular reflex with the participation of the spinal level and usually is a three-synapse reflex, which is a more objective and direct criterion in evaluation of pain and acupuncture analgesia. Given that acupuncture may inhibit the C-fiber reflex EMG, the mechanisms underlying acupuncture analgesia may be explained at both the peripheral nervous system level as well as the spinal central nervous system level.

Different sciatic nerve compression or ligation models could induce different influences on C-fiber reflexes. The CCI model can cause a diffused lesion on effenter fibers and thus damage muscle innervation of the sciatic nerve. Because damage to the motor nerve and muscular atrophy in the area innervated by the sciatic nerve may result in failure of pain behaviors and C-fiber reflex recordings (thereby affecting the evaluation of acupuncture analgesia), we aimed to establish a novel animal model to evaluate C-fiber reflex changes induced by acupuncture: namely, chronic constriction injury of the partial sciatic nerve (CCIp).

In this study, we compared the characteristics of CCIp and CCI models and utilized the CCIp model to evaluate acupuncture analgesia. In addition, we investigated the correlation between analgesia of acupuncture and the location of sciatic nerve compression.

Materials and methods

Animals

This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the Committee on the Ethics of Animal Experiments of China Academy of Chinese Medical Sciences (Approval Number: AE20110510-001). Male adult Sprague Dawley rats (200–250 g) were purchased from the Military Academy of Medical Sciences [Animal Certificate of Conformity: SCXK (Army) 2012-0004] (Beijing, People’s Republic of China). Rats were housed in clean animal facilities with the room temperature maintained at 24°C±0.5°C, humidity at 60%–70%, and noise levels lower than 60 dB. All animals were housed in groups of two or three with ad libitum access to food and water. The animals were maintained on a standard 12-hour light–dark cycle (dark cycle 8:00 PM–8:00 AM) and were allowed to acclimate to the housing conditions for 7 days prior to the experiment.

Surgical preparation

Animals were anesthetized with 10% urethane (1.0–1.2 g/kg, intraperitoneal) and placed on a sterile operating table in the left-lateral position. Before surgery, the 4-0 chromic gut (MY 41839, MYCO Medical Supplies Inc., Cary, NC, USA) was soaked in normal saline for at least 30 minutes. After shaving and sterilizing the right leg with iodophor, a transverse incision was made underneath the long head of biceps femoris and the sciatic nerve was exposed at the level of the middle of the thigh by blunt dissection for CCI of sciatic nerve trunk. And for CCIp, an oblique incision of the thigh through the skin and muscle was made and the sciatic nerve and its three terminal branches (sural, common peroneal, and tibial nerves) were exposed. Also by blunt dissection, these branches were isolated.

Four ligatures were made in both the CCI and the CCIp models. Proximal to the sciatic nerve’s trifurcation, ~7 mm of nerve was freed from the adhering tissue, and ligatures were tied around it with ~1 mm spacing in the CCI model (Figure 1A). Four ligatures were tied around the tibial nerve and the common peroneal nerve in CCIp model (Figure 1B) modified from the pSNL and SNI model. Great care was taken to avoid any damage to the sural nerve. Nerve compression was made tight enough so that toes on the fibula side twitched slightly. The innervation of the gastrocnemius was influenced by compression injury. Muscle and skin were sutured separately and carefully in layers. The neurotrophism and cutaneous sensation of the tissue were maintained as well as possible. The phenotypes of spontaneous pain and mechanical hyperalgesia after nerve ligation included hindlimb discoordination, toes curling up, and limb lameness.
Pain measurement
Mechanical pain thresholds were separately detected on the day before modeling and on the 5th and 17th day after modeling. Before pain assessment, rats were placed on the other side of a transparent partition with a stainless steel mesh bottom and allowed to move freely for 30 minutes. As modified from a previous report,19 PMWL and PMWT were measured through a Dynamic Plantar Aesthesiometer (37450, UGO, Italy) with an ascending tip which exerted a force that increased at a constant rate of 2 g/s with a cut-off force of 50 g. Stimulus was delivered to the middle of the plantar hind paw between the second and third or third and fourth metatarsals via the tip (Figure 2).20 Each hind paw was tested three times with a 15 minutes interval between each measurement, and the three measured data were averaged.

Immunohistochemistry
Animals were anesthetized with 10% urethane (1.0–1.2 g/kg, intraperitoneal) and transcardially perfused with 300 mL 4% paraformaldehyde in 0.1 M phosphate-buffered saline (PBS), pH 7.4. The L4–L6 dorsal root ganglions (DRG) were removed and placed in the same fixative (4°C) for 2 hours and then kept in 0.1 M PBS containing 25% sucrose for 48 hours. The DRG were cut into 40 μm sections on a freezing microtome. Sections were placed in 0.01 M PBS, washed three times, incubated in 0.3% Triton X-100 for 30 minutes, rinsed 3 times in 0.01 M PBS, then blocked in 10% goat serum for 1 hour. The tissue sections were incubated with a cocktail solution of primary antibodies containing calcitonin-gene-related peptide as a marker of neuro-peptidergic subpopulation (CGRP, mouse monoclonal, diluted 1:1000, ab81887, Abcam Trading (Shanghai) Company Ltd, Shanghai, People’s Republic of China), substance P as a marker of neuro-peptidergic subpopulation (SP, rabbit polyclonal, diluted 1:500, ab67006, Abcam Trading (Shanghai) Company Ltd), and Isolectin B4 as a marker of non-peptidergic sensory fibers (goat polyclonal, diluted 1:1000, sc-31339, Santa Cruz Biotechnology Inc., Dallas, TX, USA) for 15 hours at 4°C. Sections were washed three times with 0.01 M PBS and then incubated in a cocktail solution of fluorescent secondary antibodies: donkey anti-rabbit IgG-conjugated Alexa Fluor 647 (diluted 1:500, ab150075, Abcam Trading (Shanghai) Company Ltd, red fluorescence), donkey anti-mouse IgG-conjugated Alexa Fluor 488 (diluted 1:500, A21202, Invitrogen, Carlsbad, CA, USA, green fluorescence), and donkey anti-goat IgG-conjugated Alexa Fluor 405 (diluted 1:500, ab175664, Abcam Trading (Shanghai) Company Ltd, blue fluorescence) on a rocking platform shaker (protected from light) for 1 hour at room temperature. After washing three times with 0.01 M PBS, the tissue sections were mounted on glass slides, dried at room temperature protected from light for 1 hour at room temperature. After washing three times with 0.01 M PBS, the tissue sections were mounted on glass slides, dried at room temperature protected from light.
light, and sealed with coverslips. Images of the DRG slices were acquired using a confocal laser scanning microscope (LMD6000, Leica, Wetzlar, Germany) with objective magnifications of 20× and 40×. Laser channels used for excitation were 405 nm, 488 nm, and 647 nm excitation, respectively.

Electromyogram recording
Electrophysiological recordings were performed before modeling and on the 5th day after modeling. EMG of biceps femoris was recorded via a pair of noninsulated platinum–iridium electrodes. Electrical stimuli with single-square waves of 2 ms duration from Constant Current Isolated Stimulator (DS3, Digitimer Ltd., Welwyn Garden City, England) were delivered at a frequency of 0.2 Hz through stimulating electrodes which were placed in the third and fourth toes within the field of sural nerve innervation (Figure 2).13 Each stimulus was repeated 6 times. If no reflex was evoked by initial 0.5mA intensity stimulation, a stepped stimulation with 0.1mA was increased until the EMG was induced. The intervals between a set of six stimulation were 1 minute. EMG signals were amplified by Microelectrode Amplifier (SYS-DAM80, World Precision Instruments, Sarasota, FL, USA) and recorded by an acquisition system with Amplifier (SYS-DAM80, World Precision Instruments, Sarasota, FL, USA). C-fiber reflex recordings post-EA were normalized to that of the baseline pre-EA. Data with normal distributions were analyzed using the paired or unpaired t-test for two groups, and one-way or two-way analysis of variance for two or three groups. Kruskal–Wallis test was used if the data did not satisfy homogeneity of variance. P<0.05 was considered to represent statistical significance.

Results
Characteristics of CCI and CCIp rats
Animals in both the CCI and CCIp groups displayed spontaneous pain on the hind limb that underwent surgery. Symptoms of spontaneous pain were defined as hindlimb incoordination, toes curling up, and limb lameness. Three rats in CCI model group bit their own feet on the surgery side and were excluded from the study.

Pain threshold in CCI and CCIp rats
As shown in Figure 3A, no difference in PMWL was found between CCI (n=7) and CCIp (n=10) rats before modeling. After ligating the sciatic nerve in CCI or two branches of the sciatic nerve in CCIp, the PMWL decreased significantly in both groups compared to their respective pre-model measurements (CCI: 16.64±0.84 s for preligation vs 16.64±0.84 s for postligation, P<0.001; CCIp: 16.86±0.57 s for preligation vs 7.75±0.55 s for postligation, P<0.001). However, there was no significant difference between changes in both groups. As shown in Figure 3B, PMWT in CCIp rats was not significantly different from that in CCI rats before modeling (CCIp: 33.5±1.25 g; CCI: 33.08±1.72 g). PMWT in CCIp or CCI rats significantly decreased after nerve ligation (CCIp: 15.70±1.12 g; CCI: 14.35±1.20 g); however, no significant difference in PMWT was found between CCIp and CCI rats after modeling. These data suggest that ligation of either the trunk or the two branches of the sciatic nerve is capable of inducing hyperalgesia and producing neuropathic pain in the affected hind limb in rats; the phenotypes of CCIp are similar to those of CCI.
Expression of CGRP and SP in DRG

Subsequently, we observed expression of CGRP and SP in DRG of L4-6 ipsilateral compared to contralateral both in CCI and CCIp models. As shown in Figure 4A and B, CGRP and SP expression in normal rats were not apparently different between CCI and CCIp groups. After modeling, CGRP and SP expression were increased in CCI and CCIp rats. No difference was found between changes of CCI and CCIp groups (Figure 4A and B). These data suggest that ligation of the two branches of the sciatic nerve could increase the expression of CGRP and SP in a similar manner to the ligation of the trunk of the sciatic nerve.

Fiber reflex

We recorded EMG of the biceps femoris induced by Ta or Tc on the 5th day after nerve ligation. As shown in Figure 5A, EMG of the biceps femoris could be reliably induced by Ta or Tc in rats in the CCI and CCIp groups before nerve ligation. After ligating the sciatic nerve, EMG could not be triggered by Ta or Tc in CCI rats; however, in CCIp rats with sciatic branch as ligation, Tc could still induce a reliable biceps femoris EMG response while Ta failed to trigger an EMG response. As shown in Figure 5B, in CCIp (n=10) rats, C-fiber reflex threshold decreased significantly as a result of nerve branch ligation (CCIp: 0.86±0.05 mA,
pre-ligation: 1.08±0.05 mA, P<0.01). As seen from Figure 5C and D, CCIp modeling did not significantly alter the EMG latency following Tc stimulation (CCIp: 177.40±13.22 s; preligation: 152.79±6.68 s) or the duration of EMG response (CCIp: 179.24±16.64 s; preligation: 181.57±14.42 s). The above data suggest that ligation of the sciatic nerve trunk in CCI not only impaired myelinated afferent fibers but also the corresponding motor nerve, leading to dystrophy of the innervated muscles, whereas the branch ligation in CCIp did not entirely impair the sciatic motor nerve and its innervated muscles.

**EA at GB30 increased pain threshold and latency in CCI and CCIp rats**

We aimed to determine the effect of acupuncture on neuropathic mechanical pain in CCIp model. As shown in Figure 3A, in CCI (n=7) rats, EA at GB30 improved the PMWL significantly (pre-EA: 7.02±0.60 s; post-EA: 10.44±0.82 s, P<0.01); EA at GB30 also significantly increased the PMWL in CCIp (n=10) rats (pre-EA: 7.75±0.55 s; post-EA: 11.10±0.68 s, P<0.01). As can be seen from Figure 3B, in CCIp rats, EA at GB30 significantly increased PMWT (pre-EA: 15.70±1.12 g; post-EA: 22.72±1.53 g, P<0.01), similarly, a significant improvement was found in CCI rats (pre-EA: 14.35±1.20 g; post-EA: 21.35±1.60 g, P<0.001). These data suggest that EA at GB30 is capable of alleviating neuropathic mechanical pain in either CCI or CCIp rats.

**EA at GB30 decreased the C-fiber reflex EMG in CCIp rats**

To further test the effect of EA (conditioned stimulus) with different intensities at GB30 on nocuous stimulation-induced nociceptive responses, we applied EA of noxious or non-noxious intensities as 0.8 Ta, 1 Ta, 1 Tc, and 2 Tc to see the effect on 1.2 Tc-triggered C-fiber reflex EMG in normal rats (n=10) or CCIp (n=10) rats. As shown in Figure 6A, 0.8 Ta intensity of EA inhibited C-fiber reflex EMG at 0 and 1 minute in normal rats (paired t-test, **P<0.01**). Figure 6B
Model to assess acupuncture analgesia displays that 1 Ta EA significantly decreased C reflex EMG at 0, 1, 2, 3, and 4 minutes in CCIp rats but only at 0 minute in normal rats ($P<0.05$, $P<0.01$). At 3 and 4 minutes, C-fiber reflex EMG in CCIp rats was significantly lower than that in normal rats ($P<0.05$, $P<0.01$). As shown in Figure 6C, 1 Tc EA significantly reduced C-fiber reflex EMG at 0, 1, 2, and 3 minutes in CCIp rats but only at 0 and 1 minute in normal rats ($P<0.05$, $P<0.01$). As can be seen in Figure 6D, the EMG response was significantly reduced by 2 Tc EA at GB30 at 1, 2, 3, and 4 minutes in CCIp rats but at 0, 1, 2, and 3 minutes in normal rats ($P<0.05$, $P<0.01$), and the EA inhibition at 0 minute was better in normal rats than that in CCIp rats ($P<0.05$). These results suggest that EA at GB30 could alleviate nociceptive stimulation-induced EMG. The finding that EA of different intensities except 2 Tc at GB30 exhibited longer-lasting effects on 1.2 Tc-induced EMG in CCIp rats than those in normal rats implies a more significant acupuncture analgesia effect in a chronic pain model.
Discussion

In the present study, we found that ligation of the common peroneal and tibial branches of the sciatic nerve in rats was capable of inducing phenotypes of neuropathic pain and significantly decreasing PMWL and PMWT, similar to ligation of the sciatic nerve trunk. EMG of biceps femoris induced by the C-fiber reflex could be recorded on the 5th day after ligation in CCI but not CCI rats. Acupuncture at GB30 significantly increased the PMWL and PMWT in both CCI and CCIp rats. The EMG of biceps femoris triggered by the C-fiber reflex could be inhibited for more than 2 minutes by acupuncture at GB30 in CCIp rats.

Cutting the sciatic nerve axons has been established as a neuma model to mimic symptoms of phantom limb pain in humans. However, in this model, the hyperalgesia and sensory abnormalities cannot be experimentally tested. To investigate underlying mechanisms of neuropathic pain, the CCI model was established.7 Following this animal model, several alternative models were established according to different diseases mimicked. For example, the pSNL model was established by ligation of the sciatic nerve at the upper thigh level. In this model, 1/3–1/2 of the sciatic nerve is trapped in the ligature.8,13 To imitate the protrusion of lumbar intervertebral disc, the SNL model was generated by ligation of L5 spinal nerve to study pain-related emotion or pain memory.9 To imitate the nerve radicular pain caused by stenosis of the spinal canal or the intervertebral foram, a model utilizing chronic compression of the dorsal root ganglion was established.11,12 To date, CCI is still the most common research model used to study neuropathic pain in animals. CCI rats demonstrate spontaneous pain, thermal hyperalgesia, mechanical hyperalgesia, and cold hyperalgesia after nerve ligation for 1–15 days, and display peak pain expression at day 14–15. Although pain induced by nerve ligation could last for 15–30 days, hyperalgesia decreases over time in this model. Therefore, the CCI model is best utilized to mimic the symptoms and behaviors of chronic neuropathic pain in human peripheral nerve injuries (tumor suppression, heavy metal poisoning, hypoxia, or metabolic abnormalities).24 A previous study on CCI demonstrated that nerve edema could be found in the compression region 10 days after nerve ligation in rats. The edema and demyelination were exhibited in either the injury area or the distal part of ligated nerve. Generally, demyelination was found in the large myelinated axons (Aδ and Aβ), but not in the small myelinated axons (Aδ), and unmyelinated axons were not affected. Two weeks after nerve injury, the myelinated fibers including Aδ were totally lost in the distal part of the injured nerve and the unmyelinated fibers related to the proximal part decreased by 34%–84%.7 Our pain behavior data showed that both CCI and CCIp modeling could mimic neuropathic pain. However, in CCI rats, the reflex induced by Aδ or C-fiber stimulation could not be recorded 5 days after surgery, except for spontaneous EMG of the biceps femoris muscle. In CCIp rats, however, the C-fiber reflex-induced EMG could be preserved 5 days after nerve ligation. The biceps femoris of rats are dominantly innervated by L4-5 spinal nerves.25,26 Ta or Tc stimulation electrodes were performed on the third and fourth toe which are the receptive fields of the common peroneal nerve and the sural nerve. Although the damaged common peroneal nerve could impair the Aδ fiber due to common peroneal nerve demyelination in CCIp rats,7 the sural nerve, which contains Aδ fibers and the motor nerve fiber, was unaffected. We were able to record C-fiber reflex-induced EMG from this threshold stimulation on day 5 after nerve ligation in CCIp rats. On the contrary, in CCI rats, the ligation of the trunk of the sciatic nerve entirely damaged all three branches and thus led to the absence of C-fiber reflex-induced EMG.

Primary sensory neurons synthesize neuroregulator peptides such as SP and CGRP which play important roles in pain modulation.27,28 CGRP and SP are mainly expressed in small neurons with unmyelinated axons (C fibers) which are dominated by polymodal nociceptors. CGRP is also expressed in a group of medium-sized cells with finely myelinated (Aδ fibers) axons dominated by high-threshold mechanoreceptors.29 Our study shows that expression of CGRP and SP was increased in the CCIp group in a similar manner to the CCI group. Chronic demyelination and nerve injury caused impairment of the Aδ and C fibers in CCI and CCIp models and can increase the expression of these two peptides, consistent with previous reports.30,31 Our behavior and immunohistochemistry data suggest that the CCIp model may be as good as the CCI model for studying the effects and mechanisms of acupuncture analgesia on chronic neuropathic pain in experimental animals.

The C-fiber reflex is a validated biceps femoris EMG response to nociceptive stimulation at the sciatic nerve and has been commonly employed in pharmacological as well as acupuncture analgesia studies.32 As shown in nociceptive behavior and immunohistochemistry, CCIp was as good as CCI model, whereas C-fiber reflex EMG could not be induced in CCI model due to the damage of myelinated fibers of the sciatic nerve. CCIp model was preferential to CCI model because C-fiber reflex EMG can be induced in
this model to further evaluate neuropathic pain and acupuncture analgesia effects in vivo in electrophysiological studies.

GB30, a key acupoint for sciatica treatment,33 was used in this study. It is believed that deep acupuncture at GB30 may repair the damaged axons and somas by increasing the release of nerve growth factor and fructooligosaccharide.34 C-fiber reflex-induced EMG could be used to evaluate the inhibition of acupuncture on nociceptive responses. The finding that EA at GB30 could increase pain threshold as well as inhibit nociceptive reflex EMG in CCIp rats provides solid evidence that the CCIp model could be used to study acupuncture analgesia on neuropathic pain as well as its underlying mechanisms. In the future, we will further use this model to study the effects of EA of different stimulation intensities on nociceptive pain and try to elucidate the underlying biological mechanisms.

Conclusion
CCIp rats exhibited the same pain behavior and immunohistochemistry changes as CCI rats. Acupuncture at GB30 could alleviate pain behavior in both CCI and CCIp models. But C-fiber reflex EMG, which is an objective and direct criterion in evaluation of pain and acupuncture analgesia, could only be induced in CCIp model, indicating CCIp model is preferential to the CCI model for experimentally studying acupuncture analgesia on chronic neuropathic pain and its responsible mechanisms.

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Author contributions
All authors contributed toward data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.


