Effects of repeated transcranial magnetic stimulation in the dorsolateral prefrontal cortex versus motor cortex in patients with neuropathic pain after spinal cord injury: a study protocol

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ABSTRACT

Introduction Neuropathic pain is one of the common complications of spinal cord injuries (SCI), which will slow down the recovery process and result in lower quality of life. Previous studies have shown that repeated transcranial magnetic stimulation (rTMS) of the motor cortex (M1) can reduce the average pain and the most severe pain of neuropathic pain after SCI. The dorsolateral prefrontal cortex (DLPFC) area is a common target of rTMS. Recently, a few studies found that rTMS of DLPFC may relieve the neuropathic pain of SCI. So far, the efficacy of rTMS treatment in the DLPFC area in improving neuropathic pain and pain-related symptoms in patients with SCI is still unclear. Therefore, our study aims to evaluate the non-inferiority of rTMS in the DLPFC vs M1 in patients with neuropathic pain after SCI, in order to provide more options for rTMS in treating neuropathic pain after SCI.

Methods and analysis We will recruit 50 subjects with neuropathic pain after SCI. They will be randomly assigned to the DLPFC- rTMS and M1-rTMS groups and be treated with rTMS for 4 weeks. Except for the different stimulation sites, the rTMS treatment programmes of the two groups are the same: 10 Hz, 1250 pulses, 115% intensity threshold, once a day, five times a week for 4 weeks. VAS, simplified McGill Pain Questionnaire, Spinal Cord Injury Pain Date Set, Pittsburgh Sleep Quality Index and Hamilton Anxiety Scale will be evaluated at baseline, second week of treatment, fourth week of treatment and 4 weeks after the end of treatment. And VAS change will be calculated.

Ethics and dissemination The Ethics Committee of the Affiliated Hospital of Southwest Medical University has approved this trial, which is numbered KY20200041. Written informed consent will be provided to all participants after verification of the eligibility criteria. The results of the study will be published in peer-reviewed publications.

Trial registration number ChiCTR2000032362.

INTRODUCTION

Spinal cord injury (SCI) is a severe disabling disease. The annual incidence of SCI all over the world ranges from 8 to 246 per million people per year.1 One of the common complications after SCI is neuropathic pain (NPP), and about 40% of patients would suffer from persistent NPP.2 NPP is often characterised by paroxysmal, puncturing, burning, pulsating, tingling and other spontaneous or induced unpleasant abnormal sensations, which may slow down the rehabilitation process, reduce the quality of life and cause considerable social economic burden.3 8 Repeated transcranial magnetic stimulation (rTMS) is a non-invasive treatment that can cause immediate and lasting changes in cortical excitability. Clinical studies and guidelines suggested that rTMS in the motor cortex (M1) region can be used to treat NPP.5 Previous studies have shown that rTMS of M1 can reduce the average pain and the most severe pain of NPP after SCI.6 7

Strengths and limitations of this study

- This is the first clinical trial evaluating the non-inferiority of repeated transcranial magnetic stimulation (rTMS) in the dorsolateral prefrontal cortex versus M1 in patients with neuropathic pain after spinal cord injury (SCI).
- rTMS is a well-tolerated, low-risk, non-invasive intervention.
- We will make a comprehensive assessment from pain degree, pain disturbance, sleep and emotion of the patients, and provide more options for rTMS to treat neuropathic pain after SCI.
- This is a preliminary exploratory study, and the follow-up time in this trial is only 4 weeks after the end of treatment.
- This will be a single-blind study. Patients will not be blinded, so expectations about treatment may affect the accuracy of the assessment.
The dorsolateral prefrontal cortex (DLPFC) area is also a common target of rTMS for treating mood disorder and cognitive impairment, such as depression. Recent researches suggested that NPP can produce long-lasting prefrontal cortex dysfunction, manifested as pain-related cognitive dysfunction, attention shift disorder and mood disorder. The proportions of NPP with depression and anxiety are as high as 65.6% and 73.7%, respectively. Simultaneously, DLPFC area also may affect pain through several brain function networks, such as the regulation of cognitive control networks, reward/fear pathway reaction etc. It has shown that the left DLPFC area is likely to be a potential target for rTMS in treating various chronic pain diseases. A clinical trial suggests that rTMS in the DLPFC area can effectively relieve NPP in patients with SCI, but the total sample size was only 12. A study with larger sample size is needed to verify the conclusion that rTMS in the DLPFC area is effective in treating NPP. Besides, the control group in that study were sham-treated. M1 is a classic target of rTMS in the treatment of NPP. As far as we know, there is currently no randomised clinical study comparing the effects of rTMS in the DLPFC and that in M1 in the treat NPP after SCI. Considering that rTMS in DLPFC may affect multiple functions of pain, mood and cognition at the same time, we designed this non-inferiority study. Therefore, our study aims to evaluate the non-inferiority of rTMS in the DLPFC area comparing with that in M1 in patients with NPP after SCI and provide more options for rTMS to treat NPP after SCI.

METHODS

Design

The current study is a single-blind randomised controlled trial, in which patients with NPP after SCI will be randomly divided into the rTMS-DLPFC group and rTMS-M1 group (figure 1). We do not consider the sham group in the study design. Mainly because the purpose of our study is to compare the efficacy of rTMS on the DLPFC with the classical motor cortex (M1) in NPP after SCI. Besides, a previous study has shown that compared with the sham group, rTMS of the DLPFC may be effective in relieving NPP in SCI patients. Therefore, the control group in our study will be designed as rTMS of the M1 region, a positive control design, instead of a sham stimulation group. All the patients will be from the Rehabilitation Medicine Department, The Affiliated Hospital of Southwest Medical University. Importantly, physicians carrying out the assessments will not be informed of the group information.

Research objects and groups

Sample size calculation

PASS statistical software will be used to calculate the sample size. The sample size is calculated based on the Visual Analogue Scale (VAS), which is the main outcome indicator. According to previous studies, the variance of VAS after rTMS treatment in M1 area and DLPFC area is set to be 1.8. The average value of VAS after rTMS treatment in M1 area is set to be 6.0. We assume that a margin of non-inferiority of 10% is acceptable in clinical, so the margin of non-inferiority is to be 0.6. The true difference is estimated to be ~1.0. The one-sided type I error and statistical power are set to be 2.5% and 80% respectively. Each group require 21 participants at a 1:1 allocation ratio, and totally 50 participants will be needed considering a dropout rate of 15%.

Diagnostic criteria

The diagnostic criteria for SCI are as follows: (1) there is an exact cause of SCI, such as trauma or myelitis; (2) clinical manifestations conform to the definitions of quadriplegia and paraplegia in the ‘International Standard for Neurological Classification of SCI (Revised in 2011)’; (3) there are sensory and motor impairments below the injury level on the basis of the American Spinal Injury Association (ASIA) standard developed by the ASIA and (4) definitely existing SCI confirmed by CT or MR.

The diagnostic criteria for NPP are set according to the diagnostic grading standard recommended by the International Pain Research Association in 2008. The participants should meet following conditions: the pain is in a clear neuroanatomical range; the medical history suggests that there are related lesions or diseases in the peripheral or central sensory system; at least one auxiliary examination confirms Conform to the scope of neuroanatomy; at least one auxiliary examination confirms the existence of related lesions or diseases. The inclusion criteria are as follows: (1) Patients diagnosed with NPP after SCI; (2) The average VAS score of the patient self-assessment is 2.0–8.0 points in the past week; (3) Age between 18 and 60 years old; (4) At least

Figure 1 Proposed participant flow. DLPFC, dorsolateral prefrontal cortex; HAMA, Hamilton Anxiety Scale; ISICIPDS, International Spinal Cord Injury Pain Date Set; MPQ, McGill pain questionnaire; PSQI, Pittsburgh Sleep Quality Index; rTMS, repeated transcranial magnetic stimulation; VAS, Visual Analogue Scale.
The coil is held tangentially to the scalp. It achieves high-angle and distance of the stimulation coil are locked and mechanical arm through position, speed and torque. The patients must remove the metal from the head and face, such as glasses, hairpins. The figure eight coil will be placed to the DLPFC area or the M1 area according to the level of motion of each patient, the most severe pain. During the 4 weeks with treatment and 4 weeks after the treatment, all patients would receive formal treatment in the inpatient department. Once the patient would comfortably lie flat on the treatment bed. As a controller, the servo driver controls the mechanical arm through position, speed and torque. The angle and distance of the stimulation coil are locked and the coil is held tangentially to the scalp. It achieves high-precision positioning of the transmission system. The rTMS therapy will be performed by therapists who have received professional training and have been engaged in rTMS therapy for more than 3 years. The treatment room is a separate room with a quiet and safe environment. According to the level of motion of each patient, the patient would comfortably lie flat on the treatment bed. The patients must remove the metal from the head and face, such as glasses, hairpins. The figure eight coil will be placed to the DLPFC area or the M1 area according to their group. The patients are relaxed as much as possible before treatment, and the resting motor threshold (RMT) will be measured. One side of the primary motor cortex will be stimulated by a single rTMS pulse, and the motor evoked potentials will be recorded by the surface electrode at the site of stimulating the first dorsal interosseous muscle of the contralateral hand. RMT is defined as the minimum stimulation intensity that elicits a 50 μV motor evoked potential in 5 out of 10 single pulse stimulation of M1 at rest.\textsuperscript{21} 1250 pulses at an intensity of 115% RMT will be applied at 10 Hz (each train with 30 s interval). The rTMS therapy will be performed once a day, 5 days per week, and last for 4 weeks. During the study period, the participants will receive regular general care, health education and rehabilitation treatment according to their own condition, such as standing training, electrical stimulation, joint passive activity and other rehabilitation treatments. If the patient is taking anti-NPP medication regularly before participating in the trial, they do not need to stop the medication. During the trial period, if the VAS score of pain reaches 6.0 or affects sleep or rehabilitation, anti-NPP drugs can be added to the participants. During the trial, each participant’s application of drugs will be recorded in detail, including anti-NPP drugs and other drugs. If the participants have epilepsy or other conditions during the treatment, they will receive formal treatment in the inpatient department. Once the participants have epilepsy or any other conditions that prevent the patients from continuing the trial during rTMS intervention, they will drop out of the trial. The data prior to dropping out of trials will be recorded and analysed statistically.

Outcomes
Other professional evaluators will assess all patients blindly at different time points on the basis of various assessments (table 1).

In addition, we will use a questionnaire to record the baseline age, gender, symptoms, duration, ASIA grade, type of paralysis, the site of SCI, previous treatment and medication of participants (week 0). The VAS is used to evaluate the overall pain. The total length of VAS is recorded with 100 mm. The 0 point is at 0 mm, which is no pain; and the 100 points are at 100 mm, which represents the most severe pain.\textsuperscript{22} During the 4 weeks with treatment and 4 weeks after the treatment, all patients would measure the overall pain intensity by themselves using the VAS. The primary outcome is the average VAS score in the past week at each evaluation time point. We are most interested in assessment at the end of the treatment. Besides, we will calculate the VAS change at the second and fourth week of treatment, and 4 weeks after the end of treatment as it can provide additional information for pain control.

The secondary outcomes include multi-dimensional evaluation of pain, Pain interference, Sleep situation as well as Anxiety. The multidimensional evaluation of pain would be assessed using the simplified McGill Pain
related activities, interference with daily activities, mood and nocturnal sleep and (7) Is the pain being treated? The Pittsburgh Sleep Quality Index (PSQI) will be applied to evaluate the sleep quality of the participants and nocturnal sleep and (7) Is the pain being treated? The Internal Spinal Cord Injury Pain Date Set will be used to assess the degree of pain interference in daily life. Its contents include: (1) how many different types of pain there were in the past week; (2) the location of pain; (3) the type of pain; (4) the intensity of pain in the past week, assessed by the VAS; (5) the duration of each pain; (6) the effects of pain on activities, participation in recreational and social activities, satisfaction and pleasure in family-related activities, interference with daily activities, mood and nocturnal sleep and (7) Is the pain being treated? The Hamilton Anxiety Scale will be used to evaluate the anxiety situation. The Hamilton Anxiety Scale will be used for evaluation the anxiety situation. Each subitem is divided into five grades: 0, 1, 2, 3, 4. 0 means no symptoms and 4 represents severe symptoms, seriously affecting life. The total score of the scale is 56, <7 (no anxiety), 7–14 (possible anxiety), 15–20 (anxiety), 21–28 (obvious anxiety), >28 (severe anxiety).

**Evaluation time**
A total of four evaluations will be conducted, which will be evaluated before treatment, at the second and fourth week of treatment, and 4 weeks after the end of the treatment.

### Data collection and management

All recruited patients will be coded with numbers. Patient data will be collected using a case report form, including basic information, site of SCI, baseline assessment data and follow-up outcomes. The blinded physiotherapists will finish follow-up assessments when patients come back for a check. All data will be stored uniformly by the researcher, and others will not be allowed to obtain them.

### Quality assurance and safety oversight

Ethics Committee of the Affiliated Hospital of Southwest Medical University will supervise the whole process of the research.

### Statistical analysis

We will use SPSS V.23 statistical software to analyse the data and the statistical significance level will be set at α<0.05. χ² test or Fisher’s exact test will be applied to analyse the categorical data, such as gender or symptom. Data not conforming to normal distribution will be analysed using non-parametric statistical tests. Additionally, repeated measure analysis of variance and post hoc test will be used to analyse statistically significant differences in intergroup and intragroup data.

We plan to conduct the intention-to-treat analysis and per-protocol analysis at the same time in this study. Consistent results will help determine the research conclusions. If they are inconsistent, we will do further analysis and discussion. Besides, we will try our best to reduce the lack of data. If the data are missing, we will use pattern mixture models to fill the missing outcome data.

### Ethics and dissemination

The conduct of this trial will conform to the principles of the Declaration of Helsinki and ethical guidelines. This research is approved by the Ethics Committee of the Affiliated Hospital of Southwest Medical University (ethics number: KY2020041). All participants will sign the written informed consent to participate in the study. The results of the study will be published in peer-reviewed publications.
of the left DLPFC could also improve pain by enhancing brain activity of the frontal-buckle circuit involved in emotional control. These studies suggest that DLPFC is not only area activated, but it may be a critical network node related to nociceptive sensory processing and pain regulation, and also a potential intervention target for rTMS treatment. rTMS in the left prefrontal cortex can modulate the deeper limbic structures that may be participated with the affective dimension of pain, including the cingulate gyrus, hippocampus, orbitofrontal cortex and insula. Moreover, activation of the left prefrontal cortex by HF-rTMS may inhibit descending pain networks related to the periaqueductal grey and nucleus cuneiformis. rTMS has different functions depending on the frequency used. Our study plans to use 10Hz rTMS, based on previous research. Healthy adults with a 10Hz rTMS stimulation in the left prefrontal cortex demonstrated a striking increase in thermal pain thresholds. As for patients with chronic NPP, 10Hz rTMS in the DLPFC area had an average pain reduction of 19% and could increase mechanical and thermal pain thresholds significantly.

In terms of outcome evaluation, we mainly focused on pain and pain-related symptoms, including mood and sleep. There has been a lot of research to explore the relationship between them. Patients with NPP after SCI are often accompanied with different degrees of anxiety and depression. Moreover, pain and depression have a negative impact on each other through several mechanisms, and one of them may be the cognitive control. Multiple studies have shown that high-frequency DLPFC rTMS can treat depression by increasing the cognitive control of negative emotions. The DLPFC is not only a primary node within a cognitive control network but also a key network node implicated in pain modulation. Different from the motor cortex stimulation, which may directly inhibit the pain signal transmission in the spinal cord, DLPFC activation can reduce pain through cognitive control. The role of DLPFC in execution and attention is also considered to be related to the cognitive regulation of the pain process. In addition to mood disorders, about 40% of patients with NPP after SCI experience sleep disorders, which means it is difficult to fall asleep or stay asleep. Because the pain often has a two-way relationship with insomnia, with changes in one reciprocally affecting another. The study has shown that rTMS has the advantages of optimising sleep structure, improving sleep quality and maintaining therapeutic efficacy compared with drug therapy and other behavioural intervention. The rTMS on the DLPFC significantly increased the total sleep time of patients with insomnia. There is evidence that the intervention targeted at insomnia may relieve pain. Therefore, rTMS in the DLPFC area may simultaneously affect pain, sleep and mood, and establish a virtuous circle among them to replace the vicious circle. This may be the most significant advantage of the rTMS in the DLPFC area.

So, we believe that rTMS in the DLPFC may be not poorer to M1 in treating NPP with SCI. Our research is clinical research and does not involve specific neurobiological mechanisms research. The treatment of NPP...
by rTMS in the DLPCF area may be related to synaptic plasticity, some cytokines and signal pathways. For example, rTMS accommodated the brain plasticity by motivating the synthesis and release of brain-derived neurotrophic factor (BDNF) and $\gamma$-aminobutyric acid (GABA). Both BDNF and GABA were related to pain, depression and sleep regulation. If our research shows positive results, it may be necessary to strengthen neurobiological mechanisms research in the future.

There are some limitations to this trial. This is a preliminary exploratory study, so the follow-up time in this trial is only 4 weeks after the end of treatment. We will follow-up for a longer time in a future study. This will be a single-blind study. Patients will not be blinded, so expectations about treatment may affect the accuracy of the assessment. And VAS has high reliability and validity and is widely used, but it belongs to subjective pain assessment index. In addition, based on our study purpose and previous research, we do not consider the design of the sham stimulation group, so the placebo effect and the expectation of the treatment may not be completely distinguishable from the real treatment effect. And we will consider this issue in the analysis and discussion of the results after the trial is completed.

**Trial status**

This publication is based on version 1 of the rTMS protocol dated on 1 January 2021. The official start of recruitment was on 1 November 2020. The estimated end date of the trial is 1 November 2022 and recruitment of patients is ongoing.

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**Contributors**

MH and J-XW have designed this trial protocol and drafted the manuscript. J-XW and FX have gained the project funding and provided consultation during the research. CZ and Y-JX are physicians who recruit patients. XL and TW are physical therapist who using the rTMS for patients with neuropathic pain after SCI. LW and RC are responsible for the assessment at baseline and following-up. LW is a research assistant on the project who is responsible for the data collection and analysis. All authors have read and approved the final manuscript. All authors reviewed and revised the manuscript before submission and approved its content.

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**Competing interests**

None declared.

**Patient and public involvement**

Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

**Patient consent for publication**

Consent obtained directly from patient(s).

**Provenance and peer review**

Not commissioned; externally peer reviewed.

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