# Treatment effects of an electro-acupuncture therapy on stress urinary incontinence

Submission date	Recruitment status	<ul><li>Prospectively registered</li></ul>
29/06/2024	No longer recruiting	☐ Protocol
Registration date	Overall study status	Statistical analysis plan
19/07/2024	Completed	Results
Last Edited	Condition category	Individual participant data
19/07/2024	Urological and Genital Diseases	<ul><li>Record updated in last year</li></ul>

## Plain English summary of protocol

Background and study aims

This study aimed to optimise the treatment of stress urinary incontinence (SUI) using electroacupuncture and investigate the effect of electroacupuncture therapy targeted at the sacral and abdominal acupoints among patients with SUI. Stress incontinence happens when movement or activity puts pressure on the bladder, causing urine to leak. Electroacupuncture is a type of acupuncture where a small electric current is passed between pairs of acupuncture needles.

Who can participate?

Female patients aged 30-75 years with stress urinary incontinence (SUI)

What does the study involve?

Participants were randomly divided into three groups, a pelvic floor muscle training (PFMT) group, PFMT + sacral acupoint group and PFMT + sacral acupoint + abdominal acupoint group. The effects of treatment in the three groups were compared at 3 months.

What are the possible benefits and risks of participating?

SUI treatment with electroacupuncture stimulating the sacral-abdominal acupoint combined with PFMT helps patients reduce related clinical symptoms and improves their quality of life.

Where is the study run from?

Luohu Hospital of Traditional Chinese Medicine (China)

When is the study starting and how long is it expected to run for? April 2021 to January 2023

Who is funding the study?

Guangdong Provincial Bureau of Traditional Chinese Medicine (China)

Who is the main contact? Na-na Zhao, zhaonana3131@163.com

# Contact information

## Type(s)

Public, Scientific, Principal investigator

#### Contact name

Dr Na-na Zhao

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# Additional identifiers

#### Clinical Trials Information System (CTIS)

Nil known

## ClinicalTrials.gov (NCT)

Nil known

#### Protocol serial number

Nil known

# Study information

#### Scientific Title

Stress urinary incontinence treatment effect of electroacupuncture therapy targeted at sacral and abdominal acupoints in female patients

## **Study objectives**

Stress urinary incontinence (SUI) treatment with electroacupuncture stimulating the sacral-abdominal acupoint combining pelvic floor muscle training (PFMT) has certain advantages in reducing International Consultation on Incontinence Questionnaire - Short Form (ICI-Q-SF) score, reducing urinary incontinence frequency, and enhancing pelvic floor muscle strength, which helps patients to reduce related clinical symptoms and improve their quality of life.

## Ethics approval required

Ethics approval required

## Ethics approval(s)

approved 18/05/2021, Ethics Committee of Luohu Hospital of Traditional Chinese Medicine (No. 16 of Xiantong Street, Luohu District, Shenzhen, 518000, China; +86 (0)755 8231 1522; lhqzyybgs@163.com), ref: 2021051910

## Study design

Single-centre double-blinded interventional randomized controlled trial

## Primary study design

Interventional

### Study type(s)

Quality of life, Treatment

## Health condition(s) or problem(s) studied

Stress urinary incontinence (SUI)

#### **Interventions**

According to the random number table method, participants were divided into three groups: PFMT, PFMT + sacral, and PFMT + sacral + abdominal points, with 52 cases in each group.

Although the PFMT group received regular PFMT, patients in the PFMT + sacral acupoint group were treated by combining PFMT with electroacupuncture at the Huiyang, Zhonglüshu and Zhongliao acupoints in the sacral region. For patients in the PFMT + sacral acupoint + abdominal acupoint group, the Qihai, Zigong and Zhongji acupoints in the abdominal region were also involved.

Patients in the PMFT group performed the following exercises:

- 1. Kegel exercises: puerperae lay in a horizontal position breathing deeply and in a relaxed manner with flexion and the separation of both legs. During inhalation, they needed to contract the anus and urethra for 2–3 seconds without interruption. This action was repeated for 5–10 minutes once or twice a day. According to the patient's tolerance, the duration and frequency of the Kegel exercises could be adjusted and gradually increased. The longest duration of a single anus and urethra contraction was 5–10 seconds, and each exercise lasted for a maximum of 10–15 minutes.
- 2. Vaginal dumbbell training: Vaginal dumbbells suitable for the respective participants were selected based on their pelvic floor muscle strength testing results. As directed, the patients were asked to insert the vaginal dumbbell approximately 2 cm deep into the vagina and hold the dumbbell through muscle contraction. They then tried different body positions, including standing, sitting and climbing stairs (10–15 minutes at a time once a day). The above exercises were continued for 2 months.

In addition to PFMT, patients in the PFMT + sacral acupoint group underwent electroacupuncture at the Huiyang, Zhonglüshu and Zhongliao acupoints in the sacral region.

Patients in the PFMT + sacral acupoint + abdominal acupoint group also received electroacupuncture at the Qihai, Zigong and Zhongji abdominal acupoints. Specific manipulations are as follows.

The participants were asked to empty their bladders before treatment. During the course of abdominal acupuncture, patients were in a supine position, and corresponding acupuncture sites

were sterilised using routine methods. Hwato Disposable Acupuncture Needles (0.35 × 40 mm) were utilised for acupuncture in an oblique direction from the Oihai to Guanyuan acupoints for 35 mm and from the Zhongji to Qugu acupoints for 35 mm but in a straight direction at the Zigong acupoint for 25 mm. As qi is induced by acupuncture, the needle was manipulated using a uniform reinforcing-reducing method. Subsequently, the Hwato SDZ-II Electroacupuncture Apparatus was adopted to randomly connect two acupoints each time using dilatational waves, a frequency of 2/15Hz and intensity of 5 mA, all of which could be adjusted according to the patient's tolerance. The treatment was considered appropriate as long as the patients felt comfortable. This treatment continued for 30 minutes each time. During sacral acupuncture, patients were placed in a prone position, and their acupuncture sites were routinely disinfected. Hwato Disposable Acupuncture Needles  $(0.35 \times 40 \text{ mm})$  were again selected for acupuncture in an oblique direction from Huiyang to Zhonglüshu for 4–5 cm, then in a straight direction from the Zhonglüshu acupoint for 4–5 cm and obliquely upward at the Zhongliao acupoint for 4–5 cm. Through the above manipulations, a needling sensation should be continuously felt in the perineum and urinary tract region. Subsequently, an electric acupuncture device was connected. Each time, two acupoints were selected on each side of the body. The parameters configured for this device were the same as those described above. This treatment also lasted for 30 minutes, and each course of treatment was conducted once every 2 days and lasted for a period of 2 months.

## Intervention Type

Supplement

## Primary outcome(s)

Measured at 6 weeks of treatment and 3 months follow up after treatment:

- 1. Basic patient information (age, height, body mass index, educational level, past medical history and delivery mode) measured using questionnaires
- 2. 1-hour pad test results measured by pad test

# Key secondary outcome(s))

- 1. 1-hour pad test results measured by pad test at 2 weeks 4 weeks, 6 weeks of treatment and 18 weeks follow up after treatment
- 2. Times of urinary incontinence measured using International Consultation on Incontinence Questionnaire—Short Form (ICI-Q-SF) scores before treatment, during the 1-6 weeks of treatment and at 15-18 weeks follow up after treatment
- 3. Quality of life measured using Incontinence Quality of Life Instrument (I-QOL) scores before treatment, during the 1-6 weeks of treatment and at 15-18 weeks follow up after treatment 4. 72-hour urinary incontinence frequencies measured using pad test before treatment, during the 1-6 weeks of treatment and at 15-18 weeks follow up after treatment
- 5. Average pad usage per week measured using pad test before treatment, during the 1-6 weeks of treatment and at 15-18 weeks follow up after treatment

## Completion date

31/01/2023

# Eligibility

# Key inclusion criteria

- 1. Female patients aged 30–75 years
- 2. Patients with symptoms conforming to the diagnosis standards of mild and moderate SUI
- 3. Patients conforming to the syndrome differentiation standards of urinary incontinence caused

by kidney qi deficiency

4. Patients with normal routine examination results including those for urine and negative urine culture tests

## Participant type(s)

Patient

## Healthy volunteers allowed

No

## Age group

Adult

## Lower age limit

30 years

## Upper age limit

75 years

#### Sex

Female

#### Total final enrolment

156

## Key exclusion criteria

- 1. Patients with urge urinary incontinence or mixed urinary incontinence
- 2. Patients with a medical history of surgical treatment for urinary incontinence or pelvic surgery
- 3. Patients with uterine prolapse severity ≥Grade 2
- 4. Patients with symptomatic urinary system infection
- 5. Patients with residual urine volumes >30 ml
- 6. Patients with maximum uroflow rates <20 ml/s
- 7. Patients incapable of or with restricted capacity for walking, going upstairs/downstairs or running
- 8. Patients taking drugs that may affect their bladder function or receiving specific SUI treatment
- 9. Patients with SUI combined with multiple system atrophy, cauda equina lesions or myeleterosis
- 10. Patients who were pregnant or lactating
- 11. Patients fitted with cardiac pacemakers, sensitive to metals or with severe enetophobia
- 12. Patients with pelvic prolapse

## Date of first enrolment

01/06/2021

#### Date of final enrolment

31/12/2021

# Locations

#### Countries of recruitment

Study participating centre Luohu Hospital of Traditional Chinese Medicine

No. 16 Xiantong Street Luohu District Shenzhen China 518000

# Sponsor information

## Organisation

Luohu Hospital of Traditional Chinese Medicine

# Funder(s)

## Funder type

Research organisation

#### **Funder Name**

Guangdong Provincial Bureau of Traditional Chinese Medicine research project

# **Results and Publications**

## Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be available upon request from Na-na Zhao (zhaonana3131@163.com).

# IPD sharing plan summary

Available on request

# Study outputs

Output type Details Date created Date added Peer reviewed? Patient-facing?

Participant information sheet 11/11/2025 No Yes