

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

10 Jun 2026

### Effect of repetitive transcranial magnetic stimulation on dysarthria and postural disturbance in patients with progressive supranuclear palsy

#### Protocol summary

##### Study aim

we aim to measure the effect of rTMS on dysarthria and postural disturbance in patients with PSP in Mashhad on 2023

##### Design

clinical trial with intervention and sham groups, parallel, triple blind, randomized, in phase 2-3 design on 12 PSP patients, enrolled in May and Jun 2023 and followed for 4 months

##### Settings and conduct

12 PSP patients were first scored for movement (by motion kinematics analysis) and speech and swallowing (scoring by speech therapist) from patients diagnosed with PSP who were chosen from the movement disorders clinic. The place of rTMS is the psychiatry clinic, the speech and swallowing evaluation place will be the speech therapy clinic, the balance analysis place will be in the physiotherapy clinic. The patients are then randomly and blindly divided into intervention and sham groups and receive 10 sessions of rTMS every other day (in the sham group, a three cm nonconductor piece is placed between the cap of the device and the scalp) blinding of the patients and the researchers is done by sealed letters that contain codes determined based on permutation blocks.

##### Participants/Inclusion and exclusion criteria

the patients who have PSP diagnosis that have ability for walking a few steps and speaking in a few words but they don't have previous dysarthria, proprioception or vestibular sensory disorders and stroke and also who don't have rTMS contraindications

##### Intervention groups

rTMS is done by two Magstim 200 that is connected to 2 groups of flat coils and the intensity of produced MEP is until 1mV. the coil sites in contact with skull is 2 cm in laterals and 1 cm below theinion.

##### Main outcome variables

postural disturbance with motion analysis and kinematic analysis of movement and scoring based on the

calculation of the result of the measured forces; state of dysarthria and swallowing with the Pietrenberg auditory-perceptual evaluation index

#### General information

##### Reason for update

##### Acronym

##### IRCT registration information

IRCT registration number: **IRCT20230504058069N1**

Registration date: **2023-06-18, 1402/03/28**

Registration timing: **retrospective**

Last update: **2023-06-27, 1402/04/06**

Update count: **1**

##### Registration date

2023-06-18, 1402/03/28

##### Registrant information

##### Name

Mina FarahNejad

##### Name of organization / entity

##### Country

Iran (Islamic Republic of)

##### Phone

+98 912 763 5742

##### Email address

m.farahnezhad@yahoo.com

##### Recruitment status

**Recruitment complete**

##### Funding source

##### Expected recruitment start date

2023-05-26, 1402/03/05

##### Expected recruitment end date

2023-06-05, 1402/03/15

##### Actual recruitment start date

empty

##### Actual recruitment end date

empty

**Trial completion date**  
empty

**Scientific title**  
Effect of repetitive transcranial magnetic stimulation on dysarthria and postural disturbance in patients with progressive supranuclear palsy

**Public title**  
RTMS effects on PSP patients

**Purpose**  
Treatment

**Inclusion/Exclusion criteria**  
**Inclusion criteria:**  
PSP patients that was diagnosed by national institute of neurologic disease and stroke for PSP criteria patient who can stand alone and without help patient who can walk at least a few steps without help patient who can speak and answer the questions patient who has signed the informed consent form to enter the plan  
**Exclusion criteria:**  
patient who was diagnosed dementia by DSM-IV criteria patient who had vestibular or proprioception or sensory deficits patient who has any contraindication of brain stimulation (such as pacemakers or presence of metal objects in the head) the patient had dysarthria before the onset of disease patient who had past medical history of stroke patient who is mute

**Age**  
No age limit

**Gender**  
Both

**Phase**  
N/A

**Groups that have been masked**

- Participant
- Investigator
- Outcome assessor

**Sample size**  
Target sample size: 12

**Randomization (investigator's opinion)**  
Randomized

**Randomization description**  
there was used randomization techniques by blocks for patients' groups randomization, for this purpose we must make a block by 30 characters with combination of A, B alphabet in "sealed envelope" site and use it for randomization. we use allocation concealment by sealed envelopes

**Blinding (investigator's opinion)**  
Double blinded

**Blinding description**  
blinding was done in participant patients, investigators and outcome assessors. patients and investigators by using a secret code in the sealed envelopes remain blinded and information analyzer is chosen from out of researcher group Intervention patient (receive RTMS or sham) in sealed envelopes that code are received. Coding by one of the project collaborators It takes place and the doctor, evaluator and patient are blinded.

**Placebo**  
Used

**Assignment**  
Parallel

**Other design features**

**Secondary Ids**  
empty

**Ethics committees**

**1**

**Ethics committee**  
**Name of ethics committee**  
Research Ethics committees of school of medicine- Mashhad university of medical sciences  
**Street address**  
Research and Technology Vice-Chancellor, Qurashi Building, next to Hoize Cinema, University St., Mashhad  
**City**  
Mashhad  
**Province**  
Razavi Khorasan  
**Postal code**  
1496960720

**Approval date**  
2022-11-08, 1401/08/17

**Ethics committee reference number**  
IR.MUMS.MEDICAL.REC.1401.663

## Health conditions studied

**1**

**Description of health condition studied**  
PSP Progressive supranuclear palsy

**ICD-10 code**  
G23.1

**ICD-10 code description**  
Progressive supranuclear ophthalmoplegia [Steele-Richardson-Olszewski]

## Primary outcomes

**1**

**Description**  
Checking the postural instability of patients

**Timepoint**  
The beginning of the study (before starting rTMS), the end of receiving rTMS, four months later

**Method of measurement**  
Kinematic analysis of scoring movement based on the results of forces and energy

**2**

**Description**  
Checking the speech and swallowing of patients

## Timepoint

The beginning of the study (before starting rTMS), the end of receiving rTMS, four months later

## Method of measurement

Wanberg's auditory-perceptual assessment index and the Northwestern swallowing disorder test

## Secondary outcomes

### 1

#### Description

کیفیت زندگی بیماران

#### Timepoint

The beginning of the study and the end of the study

#### Method of measurement

PDQL questionnaire

## Intervention groups

### 1

#### Description

Intervention group: 6 PSP patients who receive 10 sessions of rTMS with 900 daily pulses of 10 Hz every other day. The location of the coils will be in contact with the skull two centimeters on the sides and one centimeter below the inion

#### Category

Rehabilitation

### 2

#### Description

Control group: 6 PSP patients who receive 10 sessions of rTMS every other day ,along with a 3 cm non-conductive object between the scalp and the cap of the device and 900 daily pulses of 10 Hz. The location of the coils will be in contact with the skull two centimeters on the sides and one centimeter below the inion.

#### Category

Placebo

## Recruitment centers

### 1

#### Recruitment center

##### Name of recruitment center

Clinic of movement disorders, the office of the first executive of the study

##### Full name of responsible person

Mina FarahNezhad

##### Street address

unit 303, third floor, east wing, G4 tower,g towers, before Kharazi highway,, Northern Research Blvd.,Chitgar,Tehran

##### City

Mashhad

##### Province

Razavi Khorasan

##### Postal code

1496960720

##### Phone

+98 912 763 5742

##### Email

m.farahnezhad@yahoo.com

## Sponsors / Funding sources

### 1

#### Sponsor

##### Name of organization / entity

Mashhad University of Medical Sciences

##### Full name of responsible person

dr. Ghayoor-e-mobarhan

##### Street address

Research and Technology Vice-Chancellor, Qorshi Building, next to Hoizeh Cinema, University Street,Mashhad

##### City

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##### Phone

+98 51 3841 1538

##### Email

ghayourm@mums.ac.ir

#### Grant name

#### Grant code / Reference number

#### Is the source of funding the same sponsor organization/entity?

Yes

#### Title of funding source

Mashhad University of Medical Sciences

#### Proportion provided by this source

100

#### Public or private sector

Public

#### Domestic or foreign origin

Domestic

#### Category of foreign source of funding

empty

#### Country of origin

#### Type of organization providing the funding

Academic

## Person responsible for general inquiries

#### Contact

##### Name of organization / entity

Mashhad University of Medical Sciences

##### Full name of responsible person

Mina FarahNezhad

##### Position

resident

##### Latest degree

Medical doctor

##### Other areas of specialty/work

Neurology

##### Street address

Unit 303,Eastern Wing, Third Floor,Tower G, Tower G4,before Kharazi Highway,Northern Pazhouhesh Blvd,Chitgar,Tehran

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## Person responsible for scientific inquiries

**Contact****Name of organization / entity**

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**Full name of responsible person**

mina farahnezhad

**Position**

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**Latest degree**

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## Person responsible for updating data

**Contact****Name of organization / entity**

Mashhad University of Medical Sciences

**Full name of responsible person**

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**Position**

resident

**Latest degree**

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## Sharing plan

**Deidentified Individual Participant Data Set (IPD)**

Yes - There is a plan to make this available

**Study Protocol**

Yes - There is a plan to make this available

**Statistical Analysis Plan**

Yes - There is a plan to make this available

**Informed Consent Form**

Yes - There is a plan to make this available

**Clinical Study Report**

Yes - There is a plan to make this available

**Analytic Code**

Yes - There is a plan to make this available

**Data Dictionary**

Yes - There is a plan to make this available

**Title and more details about the data/document**

All the potential data of movement score and questionnaires and speech and swallowing scores of patients can be shared after making them non-identifiable

**When the data will become available and for how long**

Access starts 6 months to a year after publication

**To whom data/document is available**

only available for people working in academic institutions

**Under which criteria data/document could be used**

It is possible to use the documents of this research only by mentioning the names of its researchers in articles and academic researches.

**From where data/document is obtainable**

from responsible author: Mina Farah Nejad Email:

m.farahnezhad@yahoo.com and

farahnezhadm991@mums.ac.ir Phone number:

00989127635742

**What processes are involved for a request to access data/document**

After requesting to receive the research documents by the researcher's academic email and after explaining about the research and the reason for the need of the desired documents by sending to the responsible author's email, if accepted, it will be sent within 1 month at the latest

**Comments**