

Clinical Trial Protocol

Iranian Registry of Clinical Trials

16 Sep 2026

Immediate effects of transcranial direct current stimulation of motor and cognitive cortex on upper extremity performance under dual-task condition in people with subacute stroke

Protocol summary

Study aim

Immediate effects of transcranial direct current stimulation of motor and cognitive cortex on upper extremity performance under dual-task condition in people with subacute stroke

Design

This study is a parallel double-blinded randomized clinical controlled trial. The referred people will be randomly allocated to one of the three groups of motor cortex stimulation, cognitive cortex stimulation and placebo

Settings and conduct

Initially, patients with stroke were referred by a neurologist based on inclusion and exclusion criteria, fulfilling a written informed consent form. Then they were randomly allocated into three groups of motor, cognition and control. After baseline assessment of outcomes, participants in motor group will receive bilateral tDCS of 1 milliampere on primary motor cortex area (M1) for 20 minutes (with cathode electrode on intact M1 and anode electrode on involved M1). Participants in cognitive group will receive bilateral tDCS of 1 milliampere on dorsolateral prefrontal cortex area (DLPFC) for 20 minutes (with cathode electrode on intact M1 and anode electrode on involved M1). Participants in control group will receive bilateral tDCS of 1 milliampere for only the initial and final 30 seconds while during the rest of the time there will be no current. The electrode placement will be similar to either of motor or cognitive group. The secondary assessment of outcomes will be performed immediately after tDCS treatment.

Participants/Inclusion and exclusion criteria

Patients with the first stroke who are at least 18 years old and have no other neurological or psychiatric problems and contraindications to the use of tDCS

Intervention groups

1.Motor cortex stimulation 2.Cognitive cortex stimulation
3.Placebo

Main outcome variables

Cognitive score, motor score, motor dual task difference, cognitive dual task difference

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20190922044839N2**

Registration date: **2021-08-22, 1400/05/31**

Registration timing: **registered_while_recruiting**

Last update: **2021-08-22, 1400/05/31**

Update count: **0**

Registration date

2021-08-22, 1400/05/31

Registrant information

Name

Tabassom Ghanavati

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 41 3337 2072

Email address

ghanavati@tbzmed.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2021-03-21, 1400/01/01

Expected recruitment end date

2021-10-23, 1400/08/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Immediate effects of transcranial direct current stimulation of motor and cognitive cortex on upper extremity performance under dual-task condition in people with subacute stroke

Public title

Immediate effects of transcranial direct current stimulation on upper extremity performance under dual-task condition in stroke

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Age over 18 years Physical ability to complete the motor activities required in examine by the examiner (ability to tap with the finger and perform a Purdepeg board test) Awareness and ability to communicate with the examiner and therapist Level of cognition enough to understand the test (score 22 or higher on the short test of mental state by the examiner) Score less than 18 upper limbs in the Fugel_Meyer test The first unilateral stroke that is at least 4 weeks old Confirmation of a stroke by one of the methods of brain imaging by a doctor

Exclusion criteria:

Contraindications to tDCS include: having a pacemaker, metal implants, a history of seizures, a skin rash, allergies or sores where the electrodes are placed, and cranial skin sensitivity. Severe upper limb pain (visual pain scale greater than 5) speech problem Existence of orthopedic or neurological disorder in the upper limb involved Upper extremity spasticity of less than 2 or more than 15 on the modified Ashworth scale

Age

From **18 years** old

Gender

Both

Phase

N/A

Groups that have been masked

- Participant
- Outcome assessor

Sample size

Target sample size: **30**

Randomization (investigator's opinion)

Randomized

Randomization description

A simple individual randomization via a statistical software will be performed. We will use the Win Pepi for making randomization order. Allocation concealment will be performed by the secretary of the assessment center

Blinding (investigator's opinion)

Double blinded

Blinding description

All participants in this study will be blind to their allocation to study groups (motor cortex stimulation, cognitive cortex stimulation and placebo). The outcome assessor, before and after intervention, will be blind to the allocation as well. However, the physiotherapist, study designer, statistical analyst and the committee of safety and data supervision will not be blind

Placebo

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

Ethics committees**1****Ethics committee****Name of ethics committee**

Ethics committee of Tabriz University of Medical Sciences

Street address

29th Bahman Blvd, Tabriz University, Faculty of Rehabilitation Sciences

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Tabriz

Province

East Azarbaijan

Postal code

5166616471

Approval date

2020-11-01, 1399/08/11

Ethics committee reference number

IR.TBZMED.REC.1399.740

Health conditions studied**1****Description of health condition studied**

Stroke

ICD-10 code

I64

ICD-10 code description

Stroke, not specified as haemorrhage or infarction

Primary outcomes**1****Description**

Cognitive test score of Auditory Stroop test

Timepoint

Before intervention and immediately after intervention

Method of measurement

Software

2

Description

Motion test score of unilateral simple Fugl_meyer test

Timepoint

Before intervention and immediately after intervention

Method of measurement

Software

3

Description

Motion test score of Purdue Pegboard board test

Timepoint

Before intervention and immediately after intervention

Method of measurement

Examiner

4

Description

Cost of cognitive dual task conditions for combining Purdue Pegboard board test and Auditory Stroop test

Timepoint

Before intervention and immediately after intervention

Method of measurement

Voice recording and calculation

5

Description

Cost of cognitive dual-task conditions for combining unilateral simple Fugl_meyer test and Auditory Stroop test

Timepoint

Before intervention and immediately after intervention

Method of measurement

Voice recording and calculation

6

Description

Cost of dual task conditions for combining Purdue Pegboard board test and Auditory Stroop test

Timepoint

Before intervention and immediately after intervention

Method of measurement

Calculation

7

Description

Cost of dual task conditions to combine unilateral simple Fugl_meyer test and Auditory Stroop test

Timepoint

Before intervention and immediately after intervention

Method of measurement

Voice recording and calculation

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group 1: Motor cortex stimulation: The participants in the motor cortex intervention group will receive one bilateral tDCS treatment session with the intensity of 1 milliamperes for 20 minutes in the primary motor area (M1). In this group, the anode electrode will be placed on the M1 area of the involved hemisphere and the cathode electrode will be placed on the M1 area of the intact hemisphere. It should be noted that the tDCS device is a nerve modulator device that includes a battery and at least two electrodes. The battery has a resistance of 9 volts. Each device must have only one anode source and one cathode evolution, one of which must be located on the head. The changes are covered with a sponge soaked in saline or other semiconductor material. The size of the changes varies between 16, 25, 35, 36 square centimeters. 2mA is the highest current strongest tested by experience, but powerful devices have increased dramatically (3-4mA). The most used current density is in the range of 0.028-0.06 current per square centimeter.

Category

Rehabilitation

2

Description

Intervention group 2: Cognitive cortex stimulation: The participants in the motor cortex intervention group will receive one bilateral tDCS treatment session with the intensity of 1 milliamperes for 20 minutes in the dorsolateral prefrontal cortex (DLPFC). In this group, the anode electrode will be placed on the DLPFC area of the involved hemisphere and the cathode electrode will be placed on the DLPFC area of the intact hemisphere. It should be noted that the tDCS device is a nerve modulator device that includes a battery and at least two electrodes. The battery has a resistance of 9 volts. Each device must have only one anode source and one cathode evolution, one of which must be located on the head. The changes are covered with a sponge soaked in saline or other semiconductor material. The size of the changes varies between 16, 25, 35, 36 square centimeters. 2mA is the highest current strongest tested by experience, but powerful devices have increased dramatically (3-4mA). The most used current density is in the range of 0.028-0.06 current per square centimeter.

Category

Rehabilitation

3

Description

Control group: Placebo: Cognitive cortex stimulation: The participants in the control group will receive a tDCS session with an electrode placement similar to either of motor intervention group or cognitive intervention group, however they will receive tDCS only for the first 30 seconds and the rest of the time will be currentless. It

should be noted that the tDCS device is a nerve modulator device that includes a battery and at least two electrodes. The battery has a resistance of 9 volts. Each device must have only one anode source and one cathode evolution, one of which must be located on the head. The changes are covered with a sponge soaked in saline or other semiconductor material. The size of the changes varies between 16, 25, 35, 36 square centimeters. 2mA is the highest current strongest tested by experience, but powerful devices have increased dramatically (3-4mA). The most used current density is in the range of 0.028-0.06 current per square centimeter.

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Physiotherapy Clinic of Tabriz Rehabilitation Faculty

Full name of responsible person

Tabassom Ghanavati

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29 Bahman Blvd, Tabriz University, Faculty of Rehabilitation Sciences

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ghanavatit@tbzmed.ac.ir

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

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No. 2 Central Building, Tabriz University of Medical Sciences, University Street

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Web page address

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Grant name

Grant code / Reference number

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

Yes

Title of funding source

Tabriz 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

Tabriz University of Medical Sciences

Full name of responsible person

Tabassom Ghanavati

Position

Assistant professor

Latest degree

Ph.D.

Other areas of specialty/work

Physiotherapy

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Person responsible for scientific inquiries

Contact

Name of organization / entity

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Full name of responsible person

Tabassom Ghanavati

Position

Assistant professor

Latest degree

Ph.D.

Other areas of specialty/work

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Person responsible for updating data**Contact****Name of organization / entity**

Tabriz University of Medical Sciences

Full name of responsible person

Tabassom Ghanavati

Position

Assistant professor

Latest degree

Ph.D.

Other areas of specialty/work

Physiotherapy

Street address

29th Bahman BLVD, Tabriz University, Rehabilitation
Faculty

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

Undecided - It is not yet known if there will be a plan to
make this available

Study Protocol

Undecided - It is not yet known if there will be a plan to
make this available

Statistical Analysis Plan

Not applicable

Informed Consent Form

Undecided - It is not yet known if there will be a plan to
make this available

Clinical Study Report

Not applicable

Analytic Code

Not applicable

Data Dictionary

Not applicable