

Clinical Trial Protocol

Iranian Registry of Clinical Trials

02 Jul 2026

Investigating the effect of binaural beats on brain electrical activity, interleukin-6 serum levels and depression severity in patients referred to the psychiatric clinic of Imam Khomeini Jiroft Hospital with major depressive disorder in 2024: A Randomized controlled trial study

Protocol summary

Study aim

Determining the effect of binaural beats on brain electrical activity, interleukin-6 serum levels, and depression severity in patients referred to the psychiatric clinic of Imam Khomeini Jiroft Hospital with major depressive disorder in 2024: A Randomized controlled trial study

Design

A controlled, four intervention groups, parallel-group, single-blind, randomized clinical trial on 96 patients. A random number table was used for randomization.

Settings and conduct

The present research will be conducted as a randomized, single-blind clinical trial in Imam Khomeini Hospital, Jiroft.

Participants/Inclusion and exclusion criteria

Inclusion criteria: Age between 20 and 60 years; having healthy hearing function; have not been exposed to binaural bits stimulation; not having any neurological disease at present or previously; not consuming alcohol; not having an underlying disease; absence of pregnancy. Exclusion criteria: lack of healthy hearing function; has ever been stimulated by binaural bits for any reason; having a neurological disease at the moment or having a history of neurological disease; alcohol consumption; having an underlying disease; being pregnant

Intervention groups

The intervention group itself is divided into 4 subgroups: a) MDD patients treated with Binaural Delta Wave, b) MDD patients treated with Binaural Theta Wave, c) MDD patients treated with Binaural Aalpha Wave, d) MDD patients treated with Binaural Gamma Wave.

Main outcome variables

Electrical brain activity, Interleukin 6 Serum level, Severity of depression, Quality of Life, sleep quality

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20230709058730N1**

Registration date: **2023-07-17, 1402/04/26**

Registration timing: **prospective**

Last update: **2023-07-17, 1402/04/26**

Update count: **0**

Registration date

2023-07-17, 1402/04/26

Registrant information

Name

Mohammad Pourfridoni

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 34 4230 6379

Email address

pourfridoni.m@jmu.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2023-08-23, 1402/06/01

Expected recruitment end date

2024-08-22, 1403/06/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Investigating the effect of binaural beats on brain electrical activity, interleukin-6 serum levels and depression severity in patients referred to the psychiatric clinic of Imam Khomeini Jiroft Hospital with major depressive disorder in 2024: A Randomized controlled trial study

Public title

Investigating the effect of binaural beats on brain electrical activity, interleukin-6 serum levels and depression severity in patients referred to the psychiatric clinic of Imam Khomeini Jiroft Hospital with major depressive disorder in 2024: A Randomized controlled trial study

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria:

All patients with major depressive disorder (MDD) who have not received any treatment so far. The age range of patients should be between 20 and 60 years. Patients must have healthy hearing function. Patients should not have been exposed to binaural beats stimulation.

Exclusion criteria:

Having or history of any neurological disease Alcohol consumption The presence of any underlying disease Pregnancy

Age

From **20 years** old to **60 years** old

Gender

Both

Phase

N/A

Groups that have been masked

- Participant

Sample size

Target sample size: **96**

Randomization (investigator's opinion)

Randomized

Randomization description

Randomization was done with a table of random numbers in such a way that the direction of reading the numbers in the table was determined from above. Numbers whose first two digits are between 00 and 12 belong to intervention group a, numbers whose first two digits are between 13 and 24 belong to intervention group b, numbers whose first two digits are between 25 and 36 belong to intervention group c, numbers which The first two digits of which are between 37 and 48 were assigned to the intervention group and the numbers whose first two digits are between 49 and 96 were assigned to the control group. Finally, one of the numbers in the table is randomly chosen and moved in the specified direction, and the numbers are recorded and assigned to study groups.

Blinding (investigator's opinion)

Single blinded

Blinding description

The present study is single-blind, so the participants (subjects) do not know which control or intervention groups they are in.

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Research Ethics Committees of Jiroft University of Medical Sciences

Street address

Pasdaran Blvd., Research and Technology Vice-Chancellor of Jiroft University of Medical Sciences

City

Jiroft

Province

Kerman

Postal code

۷۸۶۱۶۱۵۷۶۵

Approval date

2023-07-08, 1402/04/17

Ethics committee reference number

IR.JMU.REC.1402.041

Health conditions studied

1

Description of health condition studied

Major Depressive Disorder

ICD-10 code

F32

ICD-10 code description

Major depressive disorder, single episode

Primary outcomes

1

Description

Brain Electrical activity

Timepoint

The evaluation intervals of patients will be on day 0, day 14 and day 28.

Method of measurement

Electroencephalography

2

Description

Interleukin 6 Serum level

Timepoint

The evaluation intervals of patients will be on day 0, day 14 and day 28.

Method of measurement

ELISA laboratory kit

3

Description

Depression severity

Timepoint

The evaluation intervals of patients will be on day 0, day 14 and day 28.

Method of measurement

Patient Health Questionnaire (PHQ-9)

Secondary outcomes

1

Description

Quality of Life

Timepoint

The evaluation intervals of patients will be on day 0, day 14 and day 28.

Method of measurement

The World Health Organization Quality of Life - BREF (WHOQOL-BREF)

2

Description

Sleep quality

Timepoint

The evaluation intervals of patients will be on day 0, day 14 and day 28.

Method of measurement

Pittsburgh Sleep Quality Index (PSQI)

Intervention groups

1

Description

Intervention group 1: patients with major depressive disorder who are treated with binaural beat delta wave with a frequency of 1 to 3 Hz. In addition to receiving their standard medical treatment (including Escitalopram 10 mg tablet daily), the patients listen to the specified frequency for 15 minutes every afternoon for 28 days. The frequency can be played for patients through the Frequency Sound Generator (Hoel Boedec) application, which is installed by the researcher on the patient's smart phone, and patients must use headphones to hear the frequency.

Category

Treatment - Other

2

Description

Intervention group 2: patients with major depressive

disorder who are treated with binaural beat Theta wave with a frequency of 4 to 8 Hz. In addition to receiving their standard medical treatment (including Escitalopram 10 mg tablet daily), the patients listen to the specified frequency for 15 minutes every afternoon for 28 days. The frequency can be played for patients through the Frequency Sound Generator (Hoel Boedec) application, which is installed by the researcher on the patient's smart phone, and patients must use headphones to hear the frequency.

Category

Treatment - Other

3

Description

Intervention group 3: patients with major depressive disorder who are treated with binaural beat Alpha wave with a frequency of 9 to 13 Hz. In addition to receiving their standard medical treatment (including Escitalopram 10 mg tablet daily), the patients listen to the specified frequency for 15 minutes every afternoon for 28 days. The frequency can be played for patients through the Frequency Sound Generator (Hoel Boedec) application, which is installed by the researcher on the patient's smart phone, and patients must use headphones to hear the frequency.

Category

Treatment - Other

4

Description

Intervention group 4: patients with major depressive disorder who are treated with binaural beat Gamma wave with a frequency of +30 Hz. In addition to receiving their standard medical treatment (including Escitalopram 10 mg tablet daily), the patients listen to the specified frequency for 15 minutes every afternoon for 28 days. The frequency can be played for patients through the Frequency Sound Generator (Hoel Boedec) application, which is installed by the researcher on the patient's smart phone, and patients must use headphones to hear the frequency.

Category

Treatment - Other

5

Description

Control group: The control group only receives the standard medical treatment of major depressive disorder, which includes the daily consumption of one tablet of Escitalopram 10 mg.

Category

N/A

Recruitment centers

1

Recruitment center

Name of recruitment center

Imam Khomeini hospital, Jiroft
Full name of responsible person
Hedyeh Askarpour
Street address
Shafa St
City
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Province
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۷۸۶۱۶۱۵۷۶۵
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Sponsors / Funding sources

1

Sponsor

Name of organization / entity
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Full name of responsible person
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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

Jeeroft 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
Jeeroft University of Medical Sciences

Full name of responsible person

Mohammad Pourfridoni

Position

MD Student

Latest degree

A Level or less

Other areas of specialty/work

Neurology

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Name of organization / entity

Jeeroft University of Medical Sciences

Full name of responsible person

Mohammad Pourfridoni

Position

MD Student

Latest degree

A Level or less

Other areas of specialty/work

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Full name of responsible person

Mohammad Pourfridoni

Position

MD Student

Latest degree

A Level or less

Other areas of specialty/work

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Fax

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Sharing plan

Deidentified Individual Participant Data Set (IPD)

No - There is not a plan to make this available

Justification/reason for indecision/not sharing IPD

There is no further information

Study Protocol

No - There is not a plan to make this available

Statistical Analysis Plan

No - There is not a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

No - There is not a plan to make this available

Analytic Code

No - There is not a plan to make this available

Data Dictionary

No - There is not a plan to make this available