

Clinical Trial Protocol

Iranian Registry of Clinical Trials

14 Aug 2026

The effect of reflexology on post- electroconvulsive thrapy on severity of pain in depressed patients referring to educational-treatmant Razi psychiatric center of Urmia in 2018

Protocol summary

Study aim

The Effect of Reflexology on Post- Electroconvulsive thrapy on severity of Pain in depressed patients

Design

Assignment of patients to the intervention and control groups will be done in a gendered and randomized manner, and a total of 26 people will be in each group.

Settings and conduct

Control of pain intensity in patients referring to Razi Educational-treatment Psychiatric Center, Urmia, Iran

Participants/Inclusion and exclusion criteria

Inclusion criteria: age: 18-65 years old; having one of the types of depression that requires the ECT; absence of clear anomalies in the organs; conscious willingness and consent to participate in the study; not being affected by underlying diseases such as diabetes, hypothyroidism, hypoparathyroidism, musculoskeletal disorders; failure to understand perception and disturbance in clear realism; no history for anxiety disorders; no acute vision problems; earn score 3 or higher on a visual scale assessing pain; receive reflexology; no drug addiction or drug intake before the study; receive the drug at least 4 hours before the intervention. Exit criteria: no collaboration to receive intervention; illness and delirium creation in patient; cancellation of patients for continuation of study; sensitivity to touch.

Intervention groups

In the intervention group, the massage will be performed using odorless olive oil without gloves. After the initial heating of the target member, it will push the reflection points of the head and muscles in the hands and feet (middle finger and thumb) using a special stick and finally use a thumb for the solar network for 2 minutes will be pressed. All steps will be performed within 20 minutes. After 10 minutes, the severity of pain in the intervention group will be measured using the visual instrument of pain. The control group will be given

routine care.

Main outcome variables

Severity of pain

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20161116030926N1**

Registration date: **2018-09-08, 1397/06/17**

Registration timing: **prospective**

Last update: **2018-12-01, 1397/09/10**

Update count: **1**

Registration date

2018-09-08, 1397/06/17

Registrant information

Name

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 44 3275 4961

Email address

alilu@umsu.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2018-09-14, 1397/06/23

Expected recruitment end date

2018-12-14, 1397/09/23

Actual recruitment start date

empty

Actual recruitment end date

empty	number two in the control group Will take. At the end of the lottery, fourteen female patients in the intervention group and fourteen female patients in the control group as well as fourteen male patients in the intervention group and fourteen male patients in the control group will be placed.
Trial completion date	Blinding (investigator's opinion)
empty	Not blinded
Scientific title	Blinding description
The effect of reflexology on post- electroconvulsive thrapy on severity of pain in depressed patients referring to educational-treatmant Razi psychiatric center of Urmia in 2018	Placebo
Public title	Not used
The effect of reflexology on post- electroconvulsive thrapy on severity of pain in depressed patients	Assignment
Purpose	Parallel
Treatment	Other design features
Inclusion/Exclusion criteria	Secondary Ids
Inclusion criteria:	empty
Age: 18-65 years old Having one type of depression (based on documentation of the case and psychiatrist's opinion) that requires ECT Patient is not in the acute phase of psychosis Patient with obvious anomalies in the organs Patient has a willing and informed consent to participate in the study Patient with underlying diseases such as diabetes, hypothyroidism, hypoparathyroidism, musculoskeletal disorders Patient at the time of intervention with perceptual impairment and disorder The patient does not have anxiety disorders according to the diagnosis of the psychiatrist The patient has no acute visual problems. (Due to interference with the filling of the numerical scale of pain) Score 3 or higher on the visual scale of pain assessment No previous massage has been performed Drug addiction or drug intake before the study Not using the drug for at least 4 hours before the intervention	Ethics committees
Exclusion criteria:	1
The patient does not have the necessary cooperation to receive the intervention Intervention causes illness and delusions in the patient Patient disclaims continuation of the study for any reason The patient is allergic to the touch and the inability to tolerate massage in the area	Ethics committee
Age	Name of ethics committee
From 18 years old to 65 years old	Urmia University of Science & Technology Ethics Committee
Gender	Street address
Both	Urmia University of Medical Sciences,
Phase	City
N/A	Urmia
Groups that have been masked	Province
No information	West Azarbaijan
Sample size	Postal code
Target sample size: 56	5714783734
Randomization (investigator's opinion)	Approval date
Randomized	2018-08-27, 1397/06/05
Randomization description	Ethics committee reference number
Assignment of patients to intervention and control groups will be done in a gendered and randomized manner. For this purpose, in two separate and distinct bags, in sex (male or female), the number 14, number one and 14, number two, which are written on the cardboard parts. , Each patient extracts one of the numbers from the bag of his sexual group. Each patient who succeeds in making the number one in the intervention group and each patient who leaves the	IR.UMSU.REC.1397.171
	Health conditions studied
	1
	Description of health condition studied
	Severe depression without symptoms of psychosis
	ICD-10 code
	F32.2
	ICD-10 code description
	Severe depressive episode without psychotic symptoms
	Primary outcomes
	1
	Description
	Severity of pain
	Timepoint
	Measure the severity of pain before the intervention and after the intervention
	Method of measurement
	Visual Analogue Scale

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group: After initial heating, the head will press the reflection points of the head and muscles in the hands and feet (mid-thumb and middle finger) using a reflexology stick (stick) with gentle pressure and without pain, and eventually 2 minutes using the thumb to initially press the solar network at the border between the upper third and middle of the foot of the foot in the area where the foot is folded up when the foot is bent, initially fixed for 2 minutes. And then will be given a rotational massage. The pressure applied is to the extent that the upper third of the fingers of the researcher is white and the patient will feel the pressure, but will not feel pain. All of these steps will be performed within 20,

Category

Treatment - Other

2

Description

Control group: Get routine for pain control

Category

N/A

Recruitment centers

1

Recruitment center

Name of recruitment center

Educational-Treatment Razi Psychiatric Center of Urmia

Full name of responsible person

Maryam Ali ashraf jodat

Street address

Educational-Treatment Razi Psychiatric Center of Urmia Salmas Road

City

Urmia

Province

West Azarbaijan

Postal code

5716964457

Phone

+98 44 3272 2930

Fax

+98 44 3365 8921

Email

Alilu@umsu.ac.ir

Web page address

<http://razi.umsu.ac.ir>

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Oroumia University of Medical Sciences

Full name of responsible person

Dr. Iraj Mohebi

Street address

University of Science and Technology, Dept. of Research and Technology Mission Center, Emergency Department, Resalat Ave, Urmia

City

Urmia

Province

West Azarbaijan

Postal code

5714783734

Phone

+98 44 3193 7224

Fax

+98 44 3224 0642

Email

Alilu@umsu.ac.ir

Web page address

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Oroumia 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

Oroumia University of Medical Sciences

Full name of responsible person

Maryam Ali Ashraf Jodat

Position

masters student

Latest degree

Master

Other areas of specialty/work

Nursery

Street address

Faculty of Nursing and Midwifery, Nazlo, 11th Km Sero Road

City

Urmia
Province
West Azarbaijan
Postal code
5756115111
Phone
+98 44 3275 4961
Fax
+98 44 3365 8921
Email
jodat20@yahoo.com

Person responsible for scientific inquiries

Contact

Name of organization / entity
Oroumia University of Medical Sciences
Full name of responsible person
Leyla Alilu
Position
Assistant Professor
Latest degree
Ph.D.
Other areas of specialty/work
Nursery
Street address
Faculty of Nursing and Midwifery, Nazlou, 11th Km Sero Road
City
Urmia
Province
West Azarbaijan
Postal code
5756115111
Phone
+98 44 3275 4961
Fax
+98 44 3275 4921
Email
Aliluleyla@gmail.com

Person responsible for updating data

Contact

Name of organization / entity
Oroumia University of Medical Sciences
Full name of responsible person
Maryam Ali Ashraf Jodat
Position
masters student
Latest degree
Master
Other areas of specialty/work
Nursery
Street address
Faculty of Nursing and Midwifery, Nazlou, 11th Km Sero Road
City
urmia
Province
West Azarbaijan
Postal code
5756115111
Phone
+98 44 3275 4961
Fax
+98 44 3365 8921
Email
jodat20@yahoo.com

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

No - There is not a plan to make this available

Clinical Study Report

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

Analytic Code

Not applicable

Data Dictionary

Not applicable