

Clinical Trial Protocol

Iranian Registry of Clinical Trials

10 Jul 2026

The effect of auricular acupressure on the quality of life of postmenopausal women

Protocol summary

The quality of life for menopause, which in fact measures the annoying symptoms of this period

Study aim

The aim of this study is to determine the effect of auricular acupressure on the quality of life of postmenopausal women.

Design

Three arm parallel group, randomised trial, with single-blind outcome assessment

Settings and conduct

This three arm clinical trials will be conducted on postmenopausal women who have inclusion criteria. After confirming the inclusion criteria demographic and specific quality of life questionnaire for menopause will be completed. Then, the patients were randomly allocated to interventional and control groups. The sample size, including the attrition will be 45 for each group. The midwife who help to complete the questionnaires is unaware of the type of group.

Participants/Inclusion and exclusion criteria

Inclusion criteria: The patient's willingness to participate in this study Residence in Mashhad Having an age between 45-60 Non-use of tobacco Non-menstruation at least 12 months ago natural Menopause Women who have any of the physical and psychological symptoms such as flushing, sweating, sleeplessness, anxiety, depression, etc. Menopause is less than 10 years No history of surgery on the ovary No current use of any herbal medicine Not receiving hormonal and non-hormonal drugs in the last 6 months to reduce menopausal complications No history of cardiovascular disease in the past 6 months Not having thyroid disease Not having a history of abnormal vaginal bleeding Not using other complementary therapies Exclusion criteria: Unwillingness to continue cooperation Doing Acupressure less than twice a week

Intervention groups

Three groups of intervention (1-auricular acupressure with seed) and 2- control group (auricular acupressure without seed) and (routine care)

Main outcome variables

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20160731029134N2**

Registration date: **2019-01-11, 1397/10/21**

Registration timing: **registered_while_recruiting**

Last update: **2019-01-11, 1397/10/21**

Update count: **0**

Registration date

2019-01-11, 1397/10/21

Registrant information

Name

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 51 3845 1795

Email address

hadizadehz941@mums.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2018-07-23, 1397/05/01

Expected recruitment end date

2019-03-03, 1397/12/12

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

The effect of auricular acupressure on the quality of life of postmenopausal women

Public title

The effect of auricular acupressure on the quality of life of postmenopausal women

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria:

The patient's willingness to participate in this study
Residence in Mashhad Having an age between 45-60
Non-use of tobacco Non-menstruation at least 12 months ago natural Menopause Women who have any of the physical and psychological symptoms such as flushing, sweating, sleeplessness, anxiety, depression, etc
Menopause is less than 10 years No history of surgery on the ovary No current use of any herbal medicine Not receiving hormonal and non-hormonal drugs in the last 6 months to reduce menopausal complications No history of cardiovascular disease in the past 6 months Not having thyroid disease Not having a history of abnormal vaginal bleeding Not using other complementary therapies

Exclusion criteria:

Unwillingness to continue cooperation Doing Acupressure less than twice a week

Age

From **45 years** old to **60 years** old

Gender

Female

Phase

N/A

Groups that have been masked

- Outcome assessor

Sample size

Target sample size: **135**

Randomization (investigator's opinion)

Randomized

Randomization description

The eligible patients are selected using convenience sampling method and then, using random numbers table, samples will be divided into three groups A, B and C.

Blinding (investigator's opinion)

Single blinded

Blinding description

Variables are measured by the assessor (a trained midwife) who is blind to blind intervention.

Placebo

Used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics committee of Mashhad University of Medical Sciences

Street address

Avenue next to the cinema Hoveizeh- Mashhad Medical Sciences, Ghoreishi building

City

Mashhad

Province

Razavi Khorasan

Postal code

345 - 91375

Approval date

2017-05-06, 1396/02/16

Ethics committee reference number

IR.MUMS.REC.1396.23

Health conditions studied

1

Description of health condition studied

Menopausal symptoms

ICD-10 code

N95.1

ICD-10 code description

Menopausal and female climacteric states

Primary outcomes

1

Description

The quality of life for menopause, which in fact measures the annoying symptoms of this period. Determining the Four Dimensions of Quality of Life in Postmenopausal Women (Vasomotor, Psychosocial, Physical and Sexual)

Timepoint

At the beginning of the study and after four weeks

Method of measurement

Menopause-Specific Quality of Life Questionnaire

Secondary outcomes

1

Description

Signs and common complications of menopause including duration and severity of daily hot flashes

Timepoint

Daily

Method of measurement

Daily hot flash recording form

2

Description

Sleep status

Timepoint

At the beginning of the study and after four weeks

Method of measurement

Pittsburgh Sleep Quality Questionnaire

3

Description

Anxiety

Timepoint

At the beginning of the study and after four weeks

Method of measurement

Hamilton Anxiety Rating Scale

Intervention groups

1

Description

Intervention group: in the auricular acupressure group (AG), participants received ear adhesive tape with seed, on the ear sympathetic (AH6a), shenmen (TF4), adrenal gland (TG2p), subcortex (AT4), endocrine (CO18), kidney (CO10), heart (CO15) and liver (CO12) points for both ears. The patients were asked to press the acupoints by themselves four times a day for a 20 second duration each time for four weeks. It was explained that the strength of the pressing should make the local auricle congestive, flushed, hot and achy.

Category

Treatment - Other

2

Description

Control group: In one control group, participants received ear adhesive tape without seed, on the ear sympathetic (AH6a), shenmen (TF4), adrenal gland (TG2p), subcortex (AT4), endocrine (CO18), kidney (CO10), heart (CO15) and liver (CO12) points for both ears. We note control group not to touch the adhesive. To use the potential psychological effects of adhesive, people are said these adhesives can improve symptoms.

Category

Placebo

3

Description

Control group: routine care includes self-care, including doing exercise and walking, diet (such as dairy consumption, a diet containing soy, no spices) and calcium supplementation and vitamin D, exposure to sunlight wearing appropriate cotton clothing, deep breathing during flushing, avoid being near-smoker, do not smoke, recommendations to lie down, or sit on a chair and tilted her head back in the time of flushing, activities to improve sleep, such as shower before bed, eat 1 cup of warm milk, no tea and coffee at the end of

the night, ointment lubricant to reduce vaginal dryness during intercourse, it is recommended to maintain sexual relations regularly, activities to reduce anxiety such as doing yoga, and socially active.

Category

Lifestyle

Recruitment centers

1

Recruitment center

Name of recruitment center

Imam Reza Acupuncture Centers & Health Centers

Full name of responsible person

Zahra Hadizadeh-talasaz

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Mashhad University of Medical Sciences

Full name of responsible person

Mohsen Tafaghodi

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Ghoreishi building, Mashhad University of Medical Science, behind of Hoveyzeh cinema, Danesgah street, Mashhad

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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

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

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Person responsible for general inquiries

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Person responsible for updating data

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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

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

Informed Consent Form

Undecided - It is not yet known if there will be 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

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

Data Dictionary

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