

Clinical Trial Protocol

Iranian Registry of Clinical Trials

09 Jun 2026

The effect of Auriculotherapy on labor pain and Anxiety in nulliparous women

Protocol summary

Study aim

Determining the effect of auriculotherapy on the severity of pain and anxiety during vaginal delivery

Design

a clinical trial with a randomized parallel blind control group on 80 nulliparous pregnant women. This is done by the random block method

Settings and conduct

The study was performed at Shohada Hospital in Qom. In the intervention group, labels containing Vakaria seeds were affixed at 5 points in both ears, first in the dilation of 4-5 cm, then in dilatation of 7-8 cm, each point for one minute (in total 20 minutes) is pressed by the researcher. The control group affixes simple labels that are similar in color and appearance to irrelevant points and the subjects are blind and will not know the type of treatment. Severe pain with visual aids Pain and anxiety with The obvious Spielberger questionnaire is measured in the stages immediately after the intervention, dilation 7-8 cm, and the beginning of the second stage of labor.

Participants/Inclusion and exclusion criteria

Inclusion criteria: age 18-40 years, first pregnancy, gestational age 37-32 W/presentation of cephalic, BMI 26- 8/19. No wound or lesion in the corner, Cervical dilation 3-4 cm. No previous treatment with auriculotherapy, No midwife.Exclusion criteria: Lack of cooperation to continue the study, receiving epidural or spinal anesthesia, drug addiction, use of oxytocin, obstetric problems, the emergency cesarean section.

Intervention groups

In the intervention group, the researcher pasted the labels containing Vakaria seeds in 5 points of the sand, uterine, external genitalia, cerebral and autonomic in both ears, first in dilation 4-5 Cm and then in dilation 7-8 cm, each point is pressed by the researcher for one minute in the interval between uterine contractions (20 minutes in total). In the control group, simple labels that are similar in color and appearance are pasted.

Main outcome variables

Pain.Anxiety

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20171106037279N3**

Registration date: **2021-08-28, 1400/06/06**

Registration timing: **prospective**

Last update: **2021-08-28, 1400/06/06**

Update count: **0**

Registration date

2021-08-28, 1400/06/06

Registrant information

Name

farzaneh nasiri

Name of organization / entity

Shahid Beheshti University of Medical Sciences

Country

Iran (Islamic Republic of)

Phone

+98 919 453 6915

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2021-10-06, 1400/07/14

Expected recruitment end date

2022-05-25, 1401/03/04

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

The effect of Auriculotherapy on labor pain and Anxiety in nulliparous women

Public title

The effect of Auriculotherapy on labor pain and Anxiety

Purpose

Supportive

Inclusion/Exclusion criteria**Inclusion criteria:**

The mother should be 18-40 years old and the first pregnancy, the gestational age should be 37-42 weeks (term), the cephalic presentation and the pregnancy should be single, the mother should be at least literate, the mother's body mass index before pregnancy or three months The first pregnancy is 19.8-26, there is a wound or lesion in the ears, the uterine contractions spontaneously and the cervical dilatation is 3-4 cm, the mother hasn't an acute and chronic psychological problem, the mother is currently taking Related drugs. Do not be nervous, do not use cigarettes or drugs, the mother is not capable of any disease or complications of pregnancy, is not destroyed in this pregnancy, has not been treated with auricular therapy, Does not have a midwife with you.

Exclusion criteria:

Lack of cooperation to continue the study, receiving epidural or spinal anesthesia, drug addiction, use of oxytocin to induce and accelerate labor, obstetric problems during labor, emergency cesarean section before completing the study, vital signs of the mother are not in the normal range

Age

From **18 years** old to **40 years** old

Gender

Female

Phase

N/A

Groups that have been masked

- Participant

Sample size

Target sample size: **80**

Randomization (investigator's opinion)

Randomized

Randomization description

Random allocation Using the random block method (patients are randomly divided into 20 blocks of 4), research units are divided into two groups of auricular therapy, placebo. Individuals will be assigned to groups based on block randomization. Block size 4 is considered. So we have six quadruple blocks consisting of AABB, ABAB, BBAA, BABA, ABBA, BAAB. The selection of each block is random and will be done by throwing the dice twenty times (until the assignment of patients to treatment groups is completed). Assignment of treatment (auricular therapy and placebo) to the intervention and control groups will also be done randomly.

Blinding (investigator's opinion)

Single blinded

Blinding description

Subjects did not know whether they had received actual treatment or a placebo

Placebo

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Ethics committee of Qom University of Medical Sciences

Street address

Qom / Kahak section, Imam Hossain street, Reza Ahmadi house

City

Qom

Province

Ghoum

Postal code

1234567890

Approval date

2020-12-15, 1399/09/25

Ethics committee reference number

IR.MUQ.REC.1399.247

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

Pain and anxiety of nulliparous women

ICD-10 code**ICD-10 code description**

باردار

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

Pain and anxiety of nulliparous women

Timepoint

Before, during and after delivery

Method of measurement

questionnaire

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group: If the vital signs are normal, the researcher affixed the labels containing Vakaria seeds after disinfecting the ear with 70% alcohol in 5 points of sand, uterine, external genitalia, cerebral and autonomic in both ears, first in dilation of 4-5 cm and then in dilatation 7-8 cm, and each point is pressed by the researcher for one minute in the interval between uterine contractions (a total of 20 minutes)

Category

Other

2

Description

Control group: Simple tags (without Vacaria seeds) that are similar in color and appearance are pasted in 5 points: sand, uterine, external genitalia, and cerebral and autonomic in both ears.

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Shohada Hospital

Full name of responsible person

Farzaneh Nasiri

Street address

Qom / Kahak section / Imam Hossein street / Reza Ahmadi house

City

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Province

Ghous

Postal code

123456789

Phone

+98 25 3423 1395

Email

farzanehnasiri100@gmail.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Ghous University of Medical Sciences

Full name of responsible person

Farzaneh nasiri

Street address

Qom / Kahak section / Imam Hossein street / Reza Ahmadi house

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123456789

Phone

+98 25 3423 1395

Email

farzanehnasiri100@gmail.com

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Ghous 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

Ghous University of Medical Sciences

Full name of responsible person

فرزانه نصیری

Position

student

Latest degree

Master

Other areas of specialty/work

Midwifery

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Person responsible for scientific inquiries

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Name of organization / entity

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Full name of responsible person

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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

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

Title and more details about the data/document

Demographic information and information obtained from the questionnaire

When the data will become available and for how long

After publishing the article in the journal

To whom data/document is available

Researchers, Department of Medical Sciences

Under which criteria data/document could be used

Use to treat other patients

From where data/document is obtainable

Email farzanehnasiri100@ gmail.com farzaneh nasiri

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

Send message via email

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

Ghoum University of Medical Sciences

Full name of responsible person

farzaneh nasiri

Position

student

Latest degree

Master

Other areas of specialty/work

Midwifery

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