

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

05 Jun 2026

### Investigating the effect of topical *Boswellia* oil on pain severity in patients with lumbar disc herniation

#### Protocol summary

##### Study aim

Determining the effect of topical use of frankincense oil on the pain intensity of patients with lumbar disc herniation referred to neurologists' office in Rafsanjan

##### Design

The study used accessible random sampling, grouping based on pain scores and gender. Group sizes were equalized with random minimization. Sampling continued until reaching the desired size. Sample size in each group was determined to be 27.67, considering potential losses.

##### Settings and conduct

The research population was all patients with lumbar disc herniation who referred to the office of neurologists in Rafsanjan city.

##### Participants/Inclusion and exclusion criteria

Inclusion criteria: 1- Age between 20 and 60 years 2- Not suffering from nervous and mental diseases 3- Lumbar disc herniation based on neurologist and MRI diagnosis ..  
Exclusion criteria: 1- Desire to quit studying 2- The need for surgery during the study 3- The need to inject coronet during the study ..

##### Intervention groups

In the intervention group, the patients were taught to use 10 drops of frankincense oil topically twice a day for 6 weeks. In the control group, olive oil was used as a placebo in the pain area. 30 ml droppers containing olive oil, in the same packaging, for local use, were provided to patients in the placebo group.

##### Main outcome variables

Evaluation of the effect of frankincense oil on pain, level of effectiveness of frankincense oil, tolerability and side effects, relationship with images of treatment exacerbation, suggestions for prescribing the best treatment, economic effects

#### General information

##### Reason for update

##### Acronym

##### IRCT registration information

IRCT registration number: **IRCT20230919059473N1**  
Registration date: **2023-10-09, 1402/07/17**  
Registration timing: **prospective**

Last update: **2023-10-09, 1402/07/17**

Update count: **0**

##### Registration date

2023-10-09, 1402/07/17

##### Registrant information

##### Name

Fatemeh Tavakoli

##### Name of organization / entity

##### Country

Iran (Islamic Republic of)

##### Phone

+98 34 3251 8458

##### Email address

fatemetavakoli1998@gmail.com

##### Recruitment status

**Recruitment complete**

##### Funding source

##### Expected recruitment start date

2023-10-17, 1402/07/25

##### Expected recruitment end date

2024-01-15, 1402/10/25

##### Actual recruitment start date

empty

##### Actual recruitment end date

empty

##### Trial completion date

empty

##### Scientific title

Investigating the effect of topical *Boswellia* oil on pain severity in patients with lumbar disc herniation

**Public title**

effect of topical Boswellia oil on pain severity in patients with lumbar disc herniation

**Purpose**

Treatment

**Inclusion/Exclusion criteria****Inclusion criteria:**

Age between 20 and 60 years Not suffering from nervous and mental diseases Lumbar disc herniation based on neurologist and MRI diagnosis Lack of sensitivity to frankincense oil Not having wounds and lesions at the place of use Absence of diabetes mellitus Absence of absolute rest Absence of immune system problems in a person such as lupus erythematosus, multiple sclerosis (MS) Not having a history of surgery in the back area Absence of alcohol and drug addiction Not using other complementary methods for pain relief Not taking anticoagulants Absence of pregnancy and breastfeeding The amount of pain is more than 3 on the numerical-visual scale of VAS

**Exclusion criteria:**

Desire to quit studying The need for surgery during the study The need to inject coronet during the study Not performing intervention for more than two consecutive days or one alternate week Change or exacerbation of back pain Incidence of sensitivity and allergy during the implementation of the study, which is characterized by symptoms such as skin rashes.

**Age**

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

**Gender**

Both

**Phase**

1

**Groups that have been masked**

- Participant

**Sample size**

Target sample size: **60**

**Randomization (investigator's opinion)**

Randomized

**Randomization description**

The sampling method was accessible and randomly assigned to groups, classified based on important intervening variables, including pain score at the time of entering the study and gender. The number of samples in the groups was equal. In order to control the confounding factors and ensure the same number of samples in the groups, the random minimization method was used (table below). The first and second samples were randomly entered into the groups, and in the rest of the cases, attention was paid to the sum of the indicators in the groups. Sampling continued until the desired sample volume was reached.

**Blinding (investigator's opinion)**

Single blinded

**Blinding description**

The patients were completely unaware of which intervention group they were in. But they were aware of participating in the intervention.

**Placebo**

Used

**Assignment**

Parallel

**Other design features****Secondary Ids**

empty

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

Research Ethics Committees of Rafsanjan University of Medical Sciences

**Street address**

Rafsanjan University of Medical Sciences - Imam Ali Blvd

**City**

Rafsanjan

**Province**

Kerman

**Postal code**

7717933777

**Approval date**

2023-05-03, 1402/02/13

**Ethics committee reference number**

IR.RUMS.REC.1402.018

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

Patients with lumbar disc herniation

**ICD-10 code**

M51.86

**ICD-10 code description**

Other intervertebral disc disorders, lumbar region

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

Evaluation of the effect of frankincense oil on pain: This study can show whether the local consumption of frankincense oil makes a significant change in the pain intensity of lumbar disc herniation patients.

**Timepoint**

Immediately after the end of the intervention

**Method of measurement**

Visual Pain Scale Questionnaire (VAS)

**Secondary outcomes**

empty

## Intervention groups

### 1

#### Description

In the intervention group, 30 ml droppers containing 10% frankincense oil (prepared in the laboratory of the Faculty of Iranian Medicine, Kerman University of Medical Sciences) for topical use were provided to the patients. The patients were taught to use 10 drops (Razavi et al., 2019) of frankincense oil topically, twice a day for 6 weeks. It was explained to the patients that they should first massage the pain area a little and when they feel heat, put the oil on the massage area for a maximum of one minute. If the patient himself is not able to use the oil in the pain area, the second person should do it for him.

#### Category

Treatment - Other

### 2

#### Description

In the control group, olive oil was used as a placebo in the pain area. 30 ml droppers containing olive oil, in the same packaging, (prepared in the laboratory of the Faculty of Iranian Medicine, Kerman University of Medical Sciences) for topical use, were provided to patients in the placebo group. The method of use is similar to the conditions of using frankincense oil. It was emphasized to the patient to use the drops without interruption and at specified intervals. To reduce the possibility of forgetting, the patients were asked to use the oil at the same times and set a reminder on their mobile phones to remind them when to use the drops. The number of times and how the oil was consumed was tracked by the researcher once a week by phone and recorded in the relevant form.

#### Category

Treatment - Other

## Recruitment centers

### 1

#### Recruitment center

##### Name of recruitment center

Rafsanjan Health Clinic

##### Full name of responsible person

Dr. Karmi

##### Street address

Pistachio Boulevard in front of Tare Bar Square - Fars Gulf Boulevard - Rafsanjan

##### City

Rafsanjan

##### Province

Kerman

##### Postal code

0000000000

##### Phone

+98 34 3434 2215

##### Email

not\_have\_an\_email@email.com

## Sponsors / Funding sources

### 1

#### Sponsor

##### Name of organization / entity

Rafsanjan University of Medical Sciences

##### Full name of responsible person

Dr. Vahid Mohammadi Shahrokhi

##### Street address

Research and Technology Vice-Chancellor - Building No. 3 - Central Organization - Imam Ali Blvd. - Rafsanjan

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##### Province

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7717933777

##### Phone

+98 34 3428 0086

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vcrt@rums.ac.ir

#### Grant name

#### Grant code / Reference number

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

Yes

#### Title of funding source

Rafsanjan 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

Rafsanjan University of Medical Sciences

##### Full name of responsible person

Fatemeh Tavakoli

##### Position

Nursing master's student

##### Latest degree

Bachelor

##### Other areas of specialty/work

Nursery

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## Person responsible for scientific inquiries

**Contact**

**Name of organization / entity**

Rafsanjan University of Medical Sciences

**Full name of responsible person**

Dr. Shahin Heidari

**Position**

Department of Medical Surgical Nursing, School of Nursing and Midwifery, Rafsanjan University of Med

**Latest degree**

Ph.D.

**Other areas of specialty/work**

Geriatrics

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## Person responsible for updating data

**Contact**

**Name of organization / entity**

Rafsanjan University of Medical Sciences

**Full name of responsible person**

Fatemeh Tavakoli

**Position**

Nursing master's student

**Latest degree**

Bachelor

**Other areas of specialty/work**

Nursery

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**Phone**

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**Email**

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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**

Yes - There is a plan to make this available

**Statistical Analysis Plan**

Yes - There is a plan to make this available

**Informed Consent Form**

No - There is not a plan to make this available

**Clinical Study Report**

Yes - There is a plan to make this available

**Analytic Code**

Yes - There is a plan to make this available

**Data Dictionary**

No - There is not a plan to make this available

**Title and more details about the data/document**

In the analysis section, only output data and main results can be shared

**When the data will become available and for how long**

The access period starts 6 months after the results are published

**To whom data/document is available**

It will be available only for researchers working in academic and scientific institutions, or people who are also working in industry can apply to receive them.

**Under which criteria data/document could be used**

The data is available only to clarify the methodology of the present study

**From where data/document is obtainable**

Dr. Shahin Heydari - Rafsanjan University of Medical Sciences

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

An email should be sent to Dr. Shahin Heydari, and as soon as the email is seen, and at her discretion, it will be sent within 10 working days.

**Comments**