

Clinical Trial Protocol

Iranian Registry of Clinical Trials

07 Jul 2026

Comparative effectiveness of ultra-sound guided injections of corticosteroid and perineural dextrose 5% on clinical appearance, function, sonographic and electrodiagnostic findings in patients with cubital tunnel syndrome, a clinical trial study

Protocol summary

Study aim

Compare the effectiveness of ultra-sound guided injections of corticosteroid and perineural dextrose 5% on clinical appearance, function, sonographic and electrodiagnostic findings in patients with cubital tunnel syndrome

Design

The research is a randomized double blind clinical trial on 22 patients.

Settings and conduct

Samples are selected from patients with mild to moderate ulnar neuropathy referred to physical medicine and rehabilitation clinics under supervision of Shiraz University of Medical Sciences divided into two groups. Block Randomization Assignment and Double blind methods are used. After obtaining informed consent, in group A corticosteroid is injected under guide of ultrasound and in group B ultrasound guided perineural injection of dextrose 5% is done. Finally effects on clinical appearance, function, sonographic and electrodiagnostic findings are measured.

Participants/Inclusion and exclusion criteria

Inclusion criteria: Patients with age ranged from 20-65 years with diagnosis of mild to moderate ulnar neuropathy. The symptoms and signs persisted for at least 1 month. Exclusion criteria: Patients who are in a severe condition (paresis and muscle atrophy) and have brachial plexus neuropathy, history of elbow trauma or operation, rheumatologic disorder, peripheral neuropathy such as diabetes, previous perineural injection, malignancy, coagulopathy, pregnancy, active infection, or infection of injection site.

Intervention groups

Group A will receive locally 1 ml Methyl Prednisolone Acetate 40 mg and 1 ml Normal saline plus 1 ml Lidocaine 2%, under guide of ultrasound, in the cubital

tunnel. Group B will receive locally 2 ml D5W plus 1 ml Lidocaine 2% ,under guide of ultrasound, in cubital tunnel.

Main outcome variables

Pain, Sensory and motor symptoms, Activity of daily living, Cross sectional area of ulnar nerve

General information

Reason for update

There was an error in recording the date of sample collection and in calculating the dates for patient examinations and preparing the necessary conditions to start the work. The correct dates for the intervention and sampling occurred after registration in the IRCT.

Acronym

IRCT registration information

IRCT registration number: **IRCT20220929056056N1**
Registration date: **2022-12-26, 1401/10/05**
Registration timing: **prospective**

Last update: **2025-01-19, 1403/10/30**

Update count: **1**

Registration date

2022-12-26, 1401/10/05

Registrant information

Name

Maryam Mirhadi

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 71 3845 4686

Email address

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Recruitment status

Recruitment complete**Funding source****Expected recruitment start date**

2023-01-20, 1401/10/30

Expected recruitment end date

2023-08-21, 1402/05/30

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparative effectiveness of ultra-sound guided injections of corticosteroid and perineural dextrose 5% on clinical appearance, function, sonographic and electrodiagnostic findings in patients with cubital tunnel syndrome, a clinical trial study

Public title

Comparative effectiveness of ultra-sound guided injections of corticosteroid and perineural dextrose 5% in patients with cubital tunnel syndrome

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Mild to moderate ulnar nerve entrapment Age of 20-65 years At least 1 month persistence of symptoms and signs

Exclusion criteria:

Severe condition Brachial plexus neuropathy History of elbow trauma or operation Rheumatologic disorder Peripheral neuropathy Previous perineural injection Malignancy Coagulopathy Pregnancy Active infection or infection of injection site

Age

From **20 years** old to **65 years** old

Gender

Both

Phase

2

Groups that have been masked

- Care provider
- Outcome assessor
- Data analyser

Sample size

Target sample size: **22**

Randomization (investigator's opinion)

Randomized

Randomization description

Patients will be randomly allocated in to two groups by Block Randomization Assignment and Double blind methods. We will have two lists of 11 patients, including the two intervention and control groups, at random. For concealment, method of random sequencing is given to another person who is unaware of the research process, and the questionnaires are completed by a person unaware of the division of groups.

Blinding (investigator's opinion)

Double blinded

Blinding description

Participant: in this study, we does not have the ability to blind the participant because the participant is aware of receiving each intervention. Clinical care giver: we teach the caregiver how to complete the questionnaire. This person is not aware of receiving patient's intervention. Researcher: this study does not have the ability to blind the researcher due to performing both interventions by himself and being aware of receiving the kind of intervention in each group. The outcome assessor of the complete questionnaires is given to a person who is not aware of the intervention performed and he/she is asked to determine the level of performance in each person according to the questionnaires. Data analyzer: questionnaire are finally given to a person to review the information. This person does not know any of the steps of the work and the way of classification in which the intervention performed.

Placebo

Not used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Ethics committee of Shiraz University of Medical Sciences

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No 5, Yas building, 6 Alley, Ahmad Abad Square, Farhangian Town

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Province

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Postal code

7179871123

Approval date

2022-09-19, 1401/06/28

Ethics committee reference number

IR.SUMS.MED.REC.1401.318

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

Cubital tunnel syndrome

ICD-10 code

G56.20

ICD-10 code description

Lesion of ulnar nerve, unspecified upper limb

Primary outcomes

1

Description

Upper limb pain

Timepoint

Before intervention, three weeks and twelve weeks later

Method of measurement

Visual Analogue Scale, PRUNE questionnaire

2

Description

Sensory and Motor symptoms

Timepoint

Before intervention, three weeks and twelve weeks later

Method of measurement

PRUNE questionnaire

Secondary outcomes

1

Description

Function of patient

Timepoint

Before intervention, three weeks and twelve weeks later

Method of measurement

PRUNE questionnaire

2

Description

cross-sectional area of ulnar nerve

Timepoint

Before intervention and twelve weeks later

Method of measurement

Ultrasonography

Intervention groups

1

Description

First Intervention group: The patients will be placed in a supine position with the shoulder abducted and the elbow flexed at 90°.The ulnar nerve within the cubital tunnel will be identified in transverse plane, and the injection will be conducted after aseptic preparation. Patients will receive locally 1 ml Methyl Prednisolone Acetate 40 mg and 1 ml Normal saline plus 1 ml Lidocaine 2%, in one session. The patients have to complete the Visual Analog Scale and PRUNE questionnaires before entering the study and 3 and 12 weeks after intervention. Electrodiagnostic and sonographic findings will be evaluated before intervention and 12 weeks later.

Category

Rehabilitation

2

Description

Second Intervention group: The patients will be placed in a supine position with the shoulder abducted and the elbow flexed at 90°.The ulnar nerve within the cubital tunnel will be identified in transverse plane, and the injection will be conducted after aseptic preparation. Patients will receive locally 2 ml D5W plus 1 ml Lidocaine 2%, in one session. The patients have to complete the Visual Analog Scale and PRUNE questionnaires before entering the study and 3 and 12 weeks after intervention. Electrodiagnostic and sonographic findings will be evaluated before intervention and 12 weeks later.

Category

Rehabilitation

Recruitment centers

1

Recruitment center

Name of recruitment center

Imam Reza rehabilitation clinic

Full name of responsible person

Mani Ramzi

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Recruitment center

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Rajae Hospital

Full name of responsible person

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3

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Name of recruitment center
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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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7134814336
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Email
Info@sums.ac.ir
Grant name
Grant code / Reference number
Is the source of funding the same sponsor organization/entity?
No

Title of funding source
Shiraz 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
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Maryam Mirhadi
Position
Resident
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Person responsible for updating data

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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

Yes - There is a plan to make this available

Informed Consent Form

Yes - There is 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

Yes - There is a plan to make this available

Title and more details about the data/document

All available data can be shared after making people unidentifiable.

When the data will become available and for how long

Start access period one year after publishing the results.

To whom data/document is available

Everyone can access to this information.

Under which criteria data/document could be used

If the information in this study helps to improve the science process.

From where data/document is obtainable

Dr. Maryam Mirhadi, 00989177398937,
maryam.mirhadii@gmail.com

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

After sending the desired message, all authors of the study will be consulted all information will be sent within a maximum of three weeks if permitted.

Comments