

Clinical Trial Protocol

Iranian Registry of Clinical Trials

28 May 2026

Comparison of the effect of diphenhydramine with injectable lidocaine in median nerve block in patients coming to Imam Khomeini emergency department

Protocol summary

Study aim

Comparison of the effect of diphenhydramine with injectable lidocaine in median nerve block in patients coming to Imam Khomeini

Design

Clinical trial including control group, with factorial groups, triple blinded, randomised

Settings and conduct

This study is performed on patients coming to Imam Khomeini emergency department with a hand laceration. Blinding is performed in a triple blinded method. The researcher injects blindly. The project executor prepares medicines and they are put in the same syringes and they are given to the assistant with A, B and C. Independent variables are the type of anesthesia and its concentration. And dependent variables are side effects of anesthesia and the amount of pain caused by injection. The pain assessment is by using Visual analog scale. Then data will be analyzed by statistical methods.

Participants/Inclusion and exclusion criteria

Entry requirements: Patients between 16 and 65 years with hand laceration; Exit conditions: Patients over 65 and under 16 years old, Those who are allergic to diphenhydramine or lidocaine, those who receive any systemic anesthesia, those who receive MAO inhibitor

Intervention groups

A group: Control group who receive lidocaine. B group: Those who receive diphenhydramine 1%. C group: Those who receive diphenhydramine 0.5%.

Main outcome variables

Duration of medicine effect, onset of action, the amount of pain and side effects

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20190318043088N1**

Registration date: **2019-04-22, 1398/02/02**

Registration timing: **registered_while_recruiting**

Last update: **2019-04-22, 1398/02/02**

Update count: **0**

Registration date

2019-04-22, 1398/02/02

Registrant information

Name

fariba yazdanbakhsh

Name of organization / entity

Country

Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2018-07-26, 1397/05/04

Expected recruitment end date

2019-06-25, 1398/04/04

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparison of the effect of diphenhydramine with injectable lidocaine in median nerve block in patients

coming to Imam Khomeini emergency department

Public title

Comparison of the effect diphenhydramine with injectable lidocaine in median nerve block

Purpose

Supportive

Inclusion/Exclusion criteria

Inclusion criteria:

The patients coming to the emergency with hand laceration from 16 to 65 years

Exclusion criteria:

The patients under the age of 16 and over 65 People who are allergic to diphenhydramine or lidocaine or they had conractagonism for lidocaine or diphenhydramine Patients who had received any systemic pain killer patients who had received MAO inhibitor

Age

From **16 years** old to **65 years** old

Gender

Both

Phase

3

Groups that have been masked

- Participant
- Care provider
- Investigator
- Outcome assessor
- Data analyser

Sample size

Target sample size: **204**

Randomization (investigator's opinion)

Randomized

Randomization description

Selection of the sample is based on the method of the random block. Thus, the first group, A, the second group, B and the third group, C are considered based on the blocks designed and the following 25 blocks 6 randomly selected from the blocks below, all patients from the number 1 Up to 204 are coded according to one of three words A, B or C. The coding is done randomly by the supervisor of the design, and the project manager, the patient and the analyst of the results of the codes do not know anything about codes.

ABABCC/AABCBC/ABBCAC/AACBCB/CCAABB/CCBBAA/CABACB/BBAACC/CBCBAA/ABCABCCABCAB/BBCCAA/ACBACB/CABCAB/BBCACA/CAABBC/CABCBA/ACBBCA/BACBAC/AABBCC

Blinding (investigator's opinion)

Triple blinded

Blinding description

In this study, the researcher, manager, nurses and the data analyzer are kept blindness. Totally there are three groups. The first group will receive lidocaine 1%, the second group receive diphenhydramine 1% and the third group will receive diphenhydramin 0.5%. The method of blinding is triple blinding. The researcher inject medicine blindly. The medications will be prepared by the supervisor of project and taken to similar syringes and given to the assistant with A, B and C.

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics committee of Imam Khomeini hospital complex, Tehran university of Medical Sciences

Street address

Doctor Gharib Av., the end of Keshavarz Blvd.

City

Tehran

Province

Tehran

Postal code

1419733141

Approval date

2018-07-22, 1397/04/31

Ethics committee reference number

IR.TUMS.IKHC.REC.1397.095

Health conditions studied

1

Description of health condition studied

Hand laceration

ICD-10 code

S64.1

ICD-10 code description

Injury of median nerve at wrist and hand level

Primary outcomes

1

Description

Duration of medicine effect

Timepoint

After the injection of the medicine when the anesthetic begins until the anesthetic ends

Method of measurement

Chronometer

2

Description

Onset of action of medicine

Timepoint

Immediately after the injection of the medicine

Method of measurement

Chronometer

Secondary outcomes

1

Description

Side effects of medicine

Timepoint

After injection

Method of measurement

Patient Clinic and Previous Studies

2

Description

The amount of pain

Timepoint

While injecting medicine

Method of measurement

Visual analog scale

Intervention groups

1

Description

Control group: Those who receive lidocaine 1%, 3 milliliter, one time, injection, manufacturing factory Caspian

Category

Treatment - Drugs

2

Description

The first intervention group: Those who receive diphenhydramine 1%. , 3 milliliter, one time, injection, manufacturing factory Caspian

Category

Treatment - Drugs

3

Description

The second intervention group: Those who receive diphenhydramine 0.5%. , 3 milliliter, one time, injection, manufacturing factory Caspian

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Emergency department of Imam Khomeini hospital complex

Full name of responsible person

Fariba Yazdanbakhsh

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Tehran University of Medical Sciences

Full name of responsible person

Dr. Seyed Ahmad Rezaii

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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

Tehran 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

Tehran University of Medical Sciences

Full name of responsible person

Fariba Yazdanbakhsh

Position

Resident
Latest degree
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Other areas of specialty/work
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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

The total individual data of the participants will be shared after being unidentifiable.

When the data will become available and for how long

Six months after publishing results

To whom data/document is available

Doctors, dentists and researchers working in universities and academic institutes

Under which criteria data/document could be used

If someone wants to do a research related to this study, he or she can use documentation with the ethical requirements

From where data/document is obtainable

Applicants can receive documentation from the author of the article through this email: faribayazdan@yahoo.com

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

Applicants should refer to emergency department of Imam Khomeini hospital with identification card. And after confirmation, they can apply in writing. Then documentation will send them through email.

Comments