

Clinical Trial Protocol

Iranian Registry of Clinical Trials

02 Aug 2026

Comparative study of efficacy and side effects of combination Nesdonal - Fentanyl - Midazolam with Etomidate - Fentanyl - Midazolam for sedation of patients undergoing cataract surgery

Protocol summary

Study aim

Comparative study of efficacy and side effects of combination Nesdonal - Fentanyl - Midazolam with Etomidate - Fentanyl - Midazolam for sedation of patients undergoing cataract surgery at Feiz hospital in 1402

Design

This randomized clinical trial has two parallel groups, triple-blind, phase 3 on 52 patients. Random allocation software was used for randomization.

Settings and conduct

The type of study is a three-blind randomized clinical trial in the population of patients who are candidates for cataract surgery between 18 and 70 years old by phacoemulsification method in Feiz Hospital in 1401. The three-way blinding of the study: the patient, the data collector, and the data analyst do not know about injectable drugs. For the purpose of blinding, similar syringes are used without specific color and name and are covered with aluminum foil. After collecting the data until the end of the statistical analysis, one group is randomly called A group and the other B.

Participants/Inclusion and exclusion criteria

Entry criteria include: candidates for cataract surgery by emulsification method, having (ASA) between 1 and 3, age between 18 and 70 and consent to participate in the study. Conditions for not entering include: Parkinson's disease, claustrophobia, dyspnea, psychosis, acute cardiac ischemia in the last 3 months, history of narcotic drugs use, history of allergy to anesthetics, pregnancy, weighing more than 100 kg, and baseline blood pressure more than 180/110.

Intervention groups

First group: receiving Nesdonal - fentanyl - midazolam
The second group: receiving Etomidate - fentanyl - midazolam

Main outcome variables

Complications frequency; complications severity

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20230715058780N1**

Registration date: **2023-07-28, 1402/05/06**

Registration timing: **prospective**

Last update: **2023-07-28, 1402/05/06**

Update count: **0**

Registration date

2023-07-28, 1402/05/06

Registrant information

Name

SEYEDSAEED HOSSEINI

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 31 3580 3308

Email address

seyedhosseinii1377@gmail.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2023-08-05, 1402/05/14

Expected recruitment end date

2023-09-22, 1402/06/31

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparative study of efficacy and side effects of combination Nesdonal - Fentanyl - Midazolam with Etomidate - Fentanyl - Midazolam for sedation of patients undergoing cataract surgery

Public title

Comparative study of efficacy and side effects of combination Nesdonal - Fentanyl - Midazolam with Etomidate - Fentanyl - Midazolam for sedation of patients undergoing cataract surgery

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria:

Cataract surgery candidates for emulsification method Having (ASA) between 1 and 3 Age between 18 and 70 years Consent to participate in the study

Exclusion criteria:

Parkinson's Occurrence of acute cardiac ischemia in the last three months Having a history of narcotic drugs use Having a history of allergy to anesthetics Pregnancy Weighing more than one hundred kilograms Baseline blood pressure greater than 180/110 Dyspnea Psychosis

Age

From **18 years** old to **70 years** old

Gender

Both

Phase

3

Groups that have been masked

- Participant
- Outcome assessor
- Data analyser

Sample size

Target sample size: **52**

Randomization (investigator's opinion)

Randomized

Randomization description

At the beginning of each day, samples are selected based on the inclusion criteria and after informed consent, random allocation is done based on the file number by random allocation software. People are divided into two groups A and B, for the purpose of blinding, only one reliable person, other than the researcher and data collector, is aware of the type of intervention, then each group is printed in two separate but similar lists. The type of sedation and the people in each group are confirmed by respected anesthesia experts. The data collector only knows the patients by their file number and has no knowledge of the type of intervention. Also, these data are still available to the data analyst in two groups, nicknamed A and B, and the type of intervention remains with the designated person until the end of the analysis.

Blinding (investigator's opinion)

Triple blinded

Blinding description

The three-way blinding of the study in terms of: 1) the patient does not know the type of injectable drugs. 2)

The data collector has no knowledge of injectable drugs. 3) The data analyst has no knowledge of which group the data belongs to. For the purpose of blinding, similar syringes are used without a specific color and name and are covered with aluminum foil, and after collecting the data until the end of the statistical analysis, Randomly, one group is called A and the other B

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics committee of Isfahan University of Medical Sciences

Street address

Research and Technology Vice-Chancellorm., Building No. 4., Isfahan University of Medical Sciences and Health Care Services., Hazar Jarib St

City

esfahan

Province

Isfahan

Postal code

8174673461

Approval date

2023-04-01, 1402/01/12

Ethics committee reference number

IR.MUI.MED.REC.1402.015

Health conditions studied

1

Description of health condition studied

Cataract surgery

ICD-10 code

H25.0

ICD-10 code description

Age-related incipient cataract

Primary outcomes

1

Description

Blood pressure

Timepoint

before receiving the drugs, after receiving, 5, 10, 20 minutes after the start of surgery and during recovery, first immediately and then every 15 minutes until leaving the recovery

Method of measurement

Operating room monitoring device

2**Description**

Respiratory rate per minute

Timepoint

before receiving the drugs, after receiving, 5, 10, 20 minutes after the start of surgery and during recovery, first immediately and then every 15 minutes until leaving the recovery

Method of measurement

Operating room monitoring device

3**Description**

Heart rate per minute

Timepoint

before receiving the drugs, after receiving, 5, 10, 20 minutes after the start of surgery and during recovery, first immediately and then every 15 minutes until leaving the recovery

Method of measurement

Operating room monitoring device

4**Description**

Blood oxygen saturation percentage

Timepoint

before receiving the drugs, after receiving, 5, 10, 20 minutes after the start of surgery and during recovery, first immediately and then every 15 minutes until leaving the recovery

Method of measurement

Operating room monitoring device

5**Description**

Determining and comparing the average length of stay in recovery Community Verified icon

Timepoint

After transfer to recovery until the end

Method of measurement

timer

6**Description**

Determining and comparing the frequency of sedation

Timepoint

At the time of receiving the drugs and 5, 10, 20 minutes after the start of the surgery Community Verified icon

Method of measurement

Determining sedation with the Ramsay-Sedation-Scale

7**Description**

Determining and comparing the frequency of surgeon

satisfaction Community Verified icon

Timepoint

End of surgery

Method of measurement

CHECK LIST

Secondary outcomes**1****Description**

Determining and comparing the frequency of intramuscular metoclopramide consumption in recovery

Timepoint

After exiting recovery

Method of measurement

Determining nausea based on Baxter Retching Faces and metoclopramide injection in scores higher than 6 and counting doses

Intervention groups**1****Description**

Intervention group: Intervention group: The FIRST group is first injected with midazolam and fentanyl intravenous bolus with the following dose: fentanyl 1 Microg/Kg, midazolam 0.02 Mg/Kg, then after 3 minutes intravenous bolus Nesdonal is injected with the following dose: 3mg/kg

Category

Treatment - Drugs

2**Description**

Intervention group :The second group is first injected with intravenous midazolam and fentanyl bolus with the following dose: midazolam 0.02 mg/Kg, fentanyl 1 Microg/Kg, then after 3 minutes, Etomidate is injected as an infusion: 0.03mg/kg/h

Category

Treatment - Drugs

Recruitment centers**1****Recruitment center****Name of recruitment center**

Feiz Educational and Therapeutic Center

Full name of responsible person

Mitra Jabal Ameli

Street address

Feiz Educational and Therapeutic Center., Beginning of Modares St., Quds Square

City

Esfahan

Province

Isfahan

Postal code

34451020

Phone

+98 31 3445 2033

Email

feiz@mui.ac.ir

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Esfahan University of Medical Sciences

Full name of responsible person

Dr. Gholamreza Asgari

Street address

Research and Technology Vice-Chancellor., Building
No. 4., Isfahan University of Medical Sciences and
Health Care Services., Hazar Jarib St

City

Esfahan

Province

Isfahan

Postal code

8174673461

Phone

+98 31 3668 5149

Email

research@mui.ac.ir

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Esfahan 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

Esfahan University of Medical Sciences

Full name of responsible person

Dr. Mitra Jabal Ameli

Position

Professor of Anesthesiology

Latest degree

Specialist

Other areas of specialty/work

Anesthesiology

Street address

School of Medicine., Isfahan University of Medical
Sciences and Health Services., Hazar Jarib St

City

Esfahan

Province

Isfahan

Postal code

8174673461

Phone

+98 31 3620 1992

Email

jabalameli@med.mui.ac.ir

Person responsible for scientific inquiries

Contact

Name of organization / entity

Esfahan University of Medical Sciences

Full name of responsible person

Dr. Mitra Jabal Ameli

Position

Professor of Anesthesiology

Latest degree

Specialist

Other areas of specialty/work

Anesthesiology

Street address

School of Medicine., Isfahan University of Medical
Sciences and Health Services., Hazar Jarib St

City

Esfahan

Province

Isfahan

Postal code

8174673461

Phone

+98 31 3620 1992

Email

jabalameli@med.mui.ac.ir

Person responsible for updating data

Contact

Name of organization / entity

Esfahan University of Medical Sciences

Full name of responsible person

Dr. Mitra Jabal Ameli

Position

Professor of Anesthesiology

Latest degree

Specialist

Other areas of specialty/work

Anesthesiology

Street address

School of Medicine., Isfahan University of Medical
Sciences and Health Services., Hazar Jarib St

City

Esfahan

Province

Isfahan

Postal code

8174673461

Phone

+98 31 3620 1992

Email

jabalameli@med.mui.ac.ir

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