

Clinical Trial Protocol

Iranian Registry of Clinical Trials

27 Jul 2026

Evaluation of the effect of adding dexmedetomidine to del-Nido cardioplegia solution on myocardial protection in patients undergoing mitral valve replacement / repair

Protocol summary

Study aim

Evaluation of the effect of adding dexmedetomidine to Del-Nido cardioplegia solution on myocardial protection in patients undergoing mitral valve replacement/repair

Design

A controlled clinical trial with superiority, with parallel groups, three-way blind, randomized block method, phase 3 on 58 patients, www.randomization.com is used for randomization.

Settings and conduct

Patients scheduled for mitral valve repair or replacement were randomly assigned to either dexmedetomidine or placebo group. Before entering the operating room, blood samples will be sent to all patients to test for troponin I and creatinine kinase-MB. Within 6 hours of entering the Intensive Care Unit, 12 hours later and 24 hours after entering the Intensive Care Unit, the sample will be repeated and the results will be recorded. Also, patients' urinary output is recorded in the first 6 hours, the first 12 hours, and 24 hours after surgery.

Participants/Inclusion and exclusion criteria

The study population included patients seeking mitral valve repair or replacement with an ejection fraction above 40%, and no history of supraventricular dysrhythmias, no history of cardiac surgery, nephropathy following the use of contrast agents (CIN), respiratory failure, stroke and TIA, And coagulopathy.

Intervention groups

Before administration of the cardioplegia solution, a vial of 200 µg / ml dexmedetomidine (in a 2 ml syringe labeled A) per 500 ml of Del-Nido cardioplegia solution will be added to the solution. The solution will contain dexmedetomidine at a concentration of 0.4 µg / ml. The infusion of the cardioplegia solution will be continued until complete cardiac arrest with the surgeon's advice. The amount and number of times the cardioplegia solution is administered will be recorded.

Main outcome variables

Troponin and CPK-MB changes

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20210713051881N1**

Registration date: **2021-11-03, 1400/08/12**

Registration timing: **prospective**

Last update: **2021-11-03, 1400/08/12**

Update count: **0**

Registration date

2021-11-03, 1400/08/12

Registrant information

Name

Pedram Chahi

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 51 3760 3653

Email address

chahip981@mums.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2021-11-22, 1400/09/01

Expected recruitment end date

2022-05-22, 1401/03/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date
empty

Scientific title
Evaluation of the effect of adding dexmedetomidine to del-Nido cardioplegia solution on myocardial protection in patients undergoing mitral valve replacement / repair

Public title
The effect of Dexmedetomidine on myocardial protection in patients undergoing Mitral valve replacement/ repair

Purpose
Prevention

Inclusion/Exclusion criteria
Inclusion criteria:
Candidate patients for mitral valve repair or replacement with ejection fraction above 40%
Exclusion criteria:
History of supraventricular dysrhythmias History of heart surgery Nephropathy following the use of contrast agent (CIN) Persistent respiratory problems Patients with a history of Stroke and TIA Patients with a history of Coagulopathy Use an intra-aortic balloon pump (IABP) before and during surgery Use vasopressor before surgery Patient dissatisfaction to participate in the study The presence of any pathological disorder leading to the release of inflammatory cytokines Cardiopulmonary arrest before, during and after surgery

Age
From **18 years** old

Gender
Both

Phase
3

Groups that have been masked

- Participant
- Investigator
- Data and Safety Monitoring Board

Sample size
Target sample size: **56**

Randomization (investigator's opinion)
Randomized

Randomization description
Using the randomized block method and using the site www.randomization.com Block: This method is used to prevent significant imbalances in the number of participants assigned to each group. Block randomization ensures that no significant imbalance is established between groups at any time during randomization, and at certain points the number of participants in each group is equal. For this method, the volume of each block must first be determined (Example of a quadruple block). Then write a list of blocks and assign numbers to them (AABB (1) - ABAB (2) -ABBA (3) -BBAA (4) - BABA (5) - BAAB (6)) Then select random numbers between one and 6 (Eg 1 4 5 and ...) and finally specify the treatment allocation list based on previous random numbers (AB AABB-BBAA-BABA-).} Allocation Concealment Allocation Method: The lottery is done using the envelope in the package.

Blinding (investigator's opinion)
Triple blinded

Blinding description
The subjects, evaluators and analysts will be unaware of the solutions prescribed for the cardioplegia of the intervention and control groups. We will define the syringe containing the solutions with labels A and B, and the subjects, evaluators and analysts are not aware of the contents of the syringe.

Placebo
Used

Assignment
Parallel

Other design features

Secondary Ids
empty

Ethics committees

1

Ethics committee
Name of ethics committee
Ethics committee of Mashhad University of Medical Sciences
Street address
East door of the University Campus, Azadi Square, Mashhad Town 91177948064
City
Mashhad
Province
Razavi Khorasan
Postal code
91177948064

Approval date
2021-06-15, 1400/03/25

Ethics committee reference number
IR.MUMS.MEDICAL.REC.1400.193

Health conditions studied

1

Description of health condition studied
Evaluation of the effect of adding dexmedetomidine to del-Nido cardioplegia solution on myocardial protection in patients undergoing mitral valve replacement / repair

ICD-10 code
I34

ICD-10 code description
Nonrheumatic mitral valve disorders

Primary outcomes

1

Description
Troponin I Assay

Timepoint
Before entering the operating room, In the first hour, 6

hours later, 12 hours later and 24 hours after entering the Intensive Care Unit

Method of measurement

Blood sample

2

Description

Creatinine kinase _MB Assay

Timepoint

Before entering the operating room, In the first hour, 6 hours later, 12 hours later and 24 hours after entering the Intensive Care Unit

Method of measurement

Blood sample

3

Description

The rate of urinary output

Timepoint

The first 6 hours, the first 12 hours and 24 hours after the operation

Method of measurement

By catheterization

Secondary outcomes

1

Description

CRP level

Timepoint

Before surgery, 6 hours after entering the Intensive Care Unit, 12 hours after and 24 hours after entering the Intensive Care Unit

Method of measurement

Blood sample

2

Description

ESR level

Timepoint

Before surgery, 6 hours after entering the Intensive Care Unit, 12 hours after and 24 hours after entering the Intensive Care Unit

Method of measurement

Blood sample

3

Description

Inotropic Score

Timepoint

In the Intensive Care Unit

Method of measurement

Inotropic score = $([\text{dopamine} + \text{dobutamine}] \times 1) + ([\text{milrinone}] \times 15) + ([\text{epinephrine} + \text{norepinephrine} + \text{isoproterenol}] \times 100)$

Intervention groups

1

Description

Intervention group: Before administering the cardioplegia solution, a vial of 200 µg / ml dexmedetomidine (in a 2 ml syringe labeled A) per 500 ml of Del-Nido cardioplegia solution will be added to the solution. . The solution will contain dexmedetomidine at a concentration of 0.4 µg / ml. The infusion of the cardioplegia solution will be continued until complete cardiac arrest with the surgeon's advice. The amount and number of times the cardioplegia solution is administered will be recorded.

Category

Prevention

2

Description

Control group: Before administering the cardioplegia solution, 1ml of 0.9% sodium chloride solution (in a 2 ml syringe labeled B) will be added to the solution for every 500 ml of the Del-Nido cardioplegia solution. The infusion of the cardioplegia solution will be continued until complete cardiac arrest with the surgeon's advice. The amount and number of times the cardioplegia solution is administered will be recorded. The perfusionist will be unaware of the contents of the prescribed solution.

Category

Prevention

Recruitment centers

1

Recruitment center

Name of recruitment center

Imam Reza Heart Surgery Center, Mashhad, Iran

Full name of responsible person

Pedram Chahi

Street address

Imam Research and Treatment Center, Imam Reza Hospital Square, Ibne Sina St, Mashhad Town

City

Mashhad

Province

Razavi Khorasan

Postal code

9137913316

Phone

+98 51 3854 3031

Email

IRH@mums.ac.ir

2

Recruitment center

Name of recruitment center

Razavi Heart Surgery Center, Mashhad, Iran

Full name of responsible person

Pedram Chahi

Street address

Razavi Hospital, after Ghaem Bridge, Azadi Highway,
Mahhad Town

City

Mashhad

Province

Razavi Khorasan

Postal code

9198613681

Phone

+98 51 3666 8888

Email

info@razavihospital.ir

3

Recruitment center

Name of recruitment center

Shahid Rajaei Heart Surgery Center, Tehran, Iran

Full name of responsible person

Pedram Chahi

Street address

Shahid Rajaei Cardiovascular Training, Research and
Treatment Center, next to Mellat Park- corner of
Niayesh, Vali Asr St., Tehran Town

City

Tehran

Province

Tehran

Postal code

1990614331

Phone

+98 21 23921

Fax

+98 21 2204 2026

Email

NCRC@iums .ac.ir

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Mashhad University of Medical Sciences

Full name of responsible person

Majid Ghayyur Mobarhan

Street address

University St, Mashhad Town

City

Mashhad

Province

Razavi Khorasan

Postal code

1394491388

Phone

+98 51 3841 2081

Fax

+98 51 3841 3006

Email

presidentoffice@mums.ac.ir

Web page address

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Mashhad 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

Mashhad University of Medical Sciences

Full name of responsible person

Pedram Chahi

Position

Student

Latest degree

Bachelor

Other areas of specialty/work

perfusion

Street address

No. 41, Farhad Ave

City

Mashhad

Province

Razavi Khorasan

Postal code

9185745637

Phone

+98 51 3760 3653

Fax

Email

chahip981@mums.ac.ir

Person responsible for scientific inquiries

Contact

Name of organization / entity

Mashhad University of Medical Sciences

Full name of responsible person

Pedram Chahi

Position

Student

Latest degree

Bachelor

Other areas of specialty/work

perfusion

Street address

No. 41, Farhad Ave
City
Mashhad
Province
Razavi Khorasan
Postal code
9185745637
Phone
+98 51 3760 3653
Fax
Email
chahip981@mums.ac.ir

Person responsible for updating data

Contact
Name of organization / entity
Mashhad University of Medical Sciences
Full name of responsible person
Pedram Chahi
Position
Student
Latest degree
Bachelor
Other areas of specialty/work
perfusion
Street address
No. 41, Farhad Ave
City
Mashhad
Province
Razavi Khorasan
Postal code
9185745637
Phone
+98 51 3760 3653

Fax
Email
chahip981@mums.ac.ir

Sharing plan

Deidentified Individual Participant Data Set (IPD)

Yes - There is a plan to make this available

Study Protocol

No - There is not a plan to make this available

Statistical Analysis Plan

No - There is not a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

No - There is not a plan to make this available

Analytic Code

No - There is not 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

It Will be published as an article

When the data will become available and for how long

One year

To whom data/document is available

Researchers and Experts

Under which criteria data/document could be used

It would be possible after getting permission from the undersecretary of research

From where data/document is obtainable

Send email to chahip981@mums.ac.ir

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

After request, the file will be sent in Excel or Spss.

Comments