

Clinical Trial Protocol

Iranian Registry of Clinical Trials

16 Sep 2026

Evaluation of the effect of Empagliflozin on the severity of heart failure symptoms and Ecocardiography parameters in patients with congenital heart diseases with left-to-right shunt: a triple-blind, randomized, placebo-controlled trial

Protocol summary

Study aim

Improving the severity of heart failure symptoms and echocardiographic parameters in CHD patients with left-to-right shunt by administering empagliflozin

Design

A triple blind, randomised, placebo-controlled clinical trial, prospective study, superiority, parallel, phase 3, performed on 44 patients, for randomisation permuted block randomisation technique with blocks of size 6 and 2 is done using the website <https://www.sealedenvelope.com>.

Settings and conduct

44 patients with heart failure caused by congenital defects of the heart septum who are admitted to the paediatric cardiology department of Imam Reza Hospital are included in the study. Patients are randomly assigned to 2 groups intervention and control, in triple-blind manner, by assigning codes A and B to the drug and placebo groups, which only the nurse knows about the type of codes. During hospitalisation primary and secondary outcomes are evaluated.

Participants/Inclusion and exclusion criteria

Inclusion criteria: CHD patients with left-to-right shunt and heart failure aged one month to 18 years Exclusion criteria: patients with various types of cardiomyopathy myocarditi, pericarditis and cardiac tamponade thiamine deficiency severe anemia thyrotoxicosis ,pulmonary infection CKD (GFR < 20 ml/min/m2) dialysis

Intervention groups

The intervention group includes patients who receive empagliflozin tablets during hospitalization along with a standard drug regimen that includes ACEI/ARB, digoxin, and diuretics. The control group includes patients who receive placebo during hospitalization along with a standard drug regimen that includes ACEI/ARB, digoxin, and diuretics. Finally, the severity of heart failure

symptoms and echocardiography parameters in two groups are compared.

Main outcome variables

Echocardiography parameters: TRPG (tricuspid regurgitation peak gradient) Left to Right shunt gradient
Left ventricular Ejection fraction

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20120520009801N11**

Registration date: **2024-02-04, 1402/11/15**

Registration timing: **prospective**

Last update: **2024-02-04, 1402/11/15**

Update count: **0**

Registration date

2024-02-04, 1402/11/15

Registrant information

Name

Amir Hooshang Mohammadpour

Name of organization / entity

Mashhad University of Medical Sciences

Country

Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2024-03-11, 1402/12/21

Expected recruitment end date

2025-06-25, 1404/04/04

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Evaluation of the effect of Empagliflozin on the severity of heart failure symptoms and Echocardiography parameters in patients with congenital heart diseases with left-to-right shunt: a triple-blind, randomized, placebo-controlled trial

Public title

Evaluation of the effect of Empagliflozin on the severity of heart failure symptoms and echocardiography parameters in patients with congenital heart diseases with left-to-right shunt

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Patients with congenital heart diseases with left-to-right shunt Heart failure One month to 18 years

Exclusion criteria:

Patients with various types of cardiomyopathy Patients with myocarditis Patients with pericarditis and cardiac tamponade Patients with thiamine deficiency (beriberi) based on clinical symptoms Patients with severe anemia Patients with thyrotoxicosis Presence of pulmonary infection Chronic kidney disease in GFR < 20 ml/min/1.73 m² Patients on dialysis

Age

From **1 month** old to **18 years** old

Gender

Both

Phase

3

Groups that have been masked

- Participant
- Care provider
- Investigator
- Outcome assessor
- Data analyser
- Data and Safety Monitoring Board

Sample size

Target sample size: **44**

Randomization (investigator's opinion)

Randomized

Randomization description

In order to randomly assign people to the study groups while guaranteeing the equality of the sample size of the groups, permuted block randomisation technique with blocks of size 6 and 2 is done using the website <https://www.sealedenvelope.com>. For example, the allocation result could be 11 blocks of 2 with different

permutations. Type of randomization: Block Allocation Concealment: Using non-transparent sealed envelopes with a random sequence obtained from the random allocation step. Since our data will be completed over time and patients will gradually enter the study, patients are allocated to one of the control or intervention groups based on the time of entry into the study and also the sequence obtained in the randomisation stage. These codes are available to the researcher present at the hospital. It should be mentioned that this researcher is fully aware of the codes. Also, the drug or placebo is available to the researcher based on the code (A or B) and he is fully aware of which one is empagliflozin or placebo. The phone number of the researcher is also available to the patients who can call him whenever any problem arises. (This researcher is not involved in the prescription, nor in the treatment evaluation or data analysis, and is solely responsible for maintaining the codes and providing the medicine to the patients based on the random code determined by the doctor). The code assigned by him is recorded in the CRF form.

Blinding (investigator's opinion)

Triple blinded

Blinding description

After the codes were prepared and given to the researcher, if the criteria for entering the study are completed and based on the codes, the patient is randomly assigned to one of the groups. The relevant code is recorded in the CRF form and the researcher considers the medicine or placebo for the patient based on the assigned code. The patient is introduced to a nurse and the prescription of drugs is done by the nurse (therapist), meanwhile, after the patient takes the drug or placebo during the hospitalization period, the evaluator (doctor) who is different from the therapist (nurse) and does not know which medicine the patient has received and only knows the assigned code, performs the relevant evaluations. After recording, the results are given in the form of codes to the person to analyse the data and hence the data is also processed without the analyst knowing which medication is being prescribed to the patient and all information is recorded and stored confidentially without mentioning the name of the patient.

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

Vice Chancellor for Research and Technology, Third floor of Ghoreschi building, Next to Hoveyze Cinema, Daneshgah Street, Mashhad

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Mashhad

Province

Razavi Khorasan

Postal code

9138813944

Approval date

2023-10-21, 1402/07/29

Ethics committee reference number

IR.MUMS.REC.1402.229

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

Congenital heart disease with left to right shunt causing heart failure symptoms

ICD-10 code

Q21

ICD-10 code description

Congenital malformations of cardiac septa

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

TRPG (tricuspid regurgitation peak gradient)

Timepoint

At the beginning and at the end of hospitalization

Method of measurement

Echocardiography

2**Description**

Left to Right shunt gradient

Timepoint

At the beginning and at the end of hospitalization

Method of measurement

Echocardiography

3**Description**

Left ventricular Ejection fraction

Timepoint

At the beginning and at the end of hospitalization

Method of measurement

Echocardiography

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

Heart failure symptoms

Timepoint

Daily

Method of measurement

Age-based Ross classification for heart failure in children

2**Description**

Vital signs

Timepoint

Daily

Method of measurement

Sphygmomanometer, Thermometer, Vital sign monitoring device , Nurse

3**Description**

Oxygen saturation (Spo2)

Timepoint

Daily

Method of measurement

Pulse oximeter

4**Description**

Weight

Timepoint

Daily

Method of measurement

Scale

5**Description**

Inflammatory markers:CRP and ESR

Timepoint

At the beginning and at the end of hospitalisation

Method of measurement

laboratory tests

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

Intervention group: patients who receive empagliflozin tablets during hospitalization along with a standard drug regimen that includes ACEI/ARB, digoxin, and diuretics

Category

Treatment - Drugs

2**Description**

Control group: patients who receive placebo during hospitalization along with a standard drug regimen that includes ACEI/ARB, digoxin, and diuretics.

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Paediatric cardiology department of Imam Reza hospital

Full name of responsible person

Amirhooshang Mohamadpour

Street address

Imam Reza hospital, Imam Reza Square, Ibn-e Sina. Avenue, Mashhad , Iran. City. Mashhad. Province. Razavi Khorasan.

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Mashhad University of Medical Sciences

Full name of responsible person

Mohsen Tafaghodi

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Web page address

<https://v-research.mums.ac.ir/moavenat/2021-04-06-05-57-55>

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

Amir Hooshang Mohammadpour

Position

Professor

Latest degree

Ph.D.

Other areas of specialty/work

Medical Pharmacy

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Person responsible for scientific inquiries

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Position

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Sharing plan**Deidentified Individual Participant Data Set (IPD)**

Yes - There is 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

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 data is potentially shareable after unidentifiable individuals

When the data will become available and for how long

Six months after the results are published

To whom data/document is available

People working in academic institutions

Under which criteria data/document could be used

Use in clinic to improve disease severity, In the condition that safety and efficacy are confirmed in the study

From where data/document is obtainable

Dr amirhoushang mohamadpour Email:
Mohamadpoorah@mums.ac.ir

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

Reply to email within a maximum of one week

Comments