

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

18 Jul 2026

### Evaluating the effect of empagliflozin on blood pressure, cardiovascular events, mortality and renal parameters in patients undergoing coronary artery bypass surgery admitted to the A coronary care unit

#### Protocol summary

##### Study aim

Evaluating the effect of empagliflozin on blood pressure, cardiovascular events, mortality and renal parameters in patients undergoing coronary artery bypass surgery

##### Design

Double-blind, randomized, placebo-controlled clinical

##### Settings and conduct

This study will be conducted in the intensive care unit of Afshar Hospital in Yazd. The patients will be divided into two groups receiving Empagliflozin tablet and placebo by random permutation block method. Each patient will be identified with a number and the list of numbers of people who should be placed in each group will be given to the nurses and the participants, researchers, doctors And the data collectors will be unaware of this list.

##### Participants/Inclusion and exclusion criteria

Input criteria include: Patients undergoing coronary artery bypass surgery Not being allergic to empagliflozin  
Exclusion criteria include: Patients who cannot get enough information with echocardiography History of taking pioglitazone at least 8 weeks before entering the study Glomerular filtration rate less than 30 mg per minute per 1.73 square meters of body surface based on the CKD-EPI formula Pregnant or lactating women

##### Intervention groups

intervention group: Recipients of empagliflozin 10 mg tablets from Abidi Pharmaceutical Company receive one empagliflozin tablet orally a few hours after coronary artery bypass surgery to reduce blood pressure, cardiovascular events, mortality, and control kidney parameters control group: The recipients of a placebo prepared in the pharmaceutical laboratory of the Yazd Faculty of Pharmacy in the same size and color as the empagliflozin tablet, a few hours after coronary bypass surgery, take a 10 mg tablet orally to reduce blood pressure, cardiovascular events, mortality and Renal controls

##### Main outcome variables

The rate of reduction of blood pressure, cardiovascular events, mortality and control of renal parameters

#### General information

##### Reason for update

##### Acronym

##### IRCT registration information

IRCT registration number: **IRCT20190911044744N3**

Registration date: **2023-12-03, 1402/09/12**

Registration timing: **prospective**

Last update: **2023-12-03, 1402/09/12**

Update count: **0**

##### Registration date

2023-12-03, 1402/09/12

##### Registrant information

##### Name

Ehsan Mirzaei

##### Name of organization / entity

##### Country

Iran (Islamic Republic of)

##### Phone

+98 35 1820 5885

##### Email address

ehsan.mirzaei.1369@gmail.com

##### Recruitment status

##### Recruitment complete

##### Funding source

##### Expected recruitment start date

2024-01-01, 1402/10/11

##### Expected recruitment end date

2024-06-30, 1403/04/10

##### Actual recruitment start date

empty

**Actual recruitment end date**

empty

**Trial completion date**

empty

**Scientific title**

Evaluating the effect of empagliflozin on blood pressure, cardiovascular events, mortality and renal parameters in patients undergoing coronary artery bypass surgery admitted to the A coronary care unit

**Public title**

Effect of empagliflozin on blood pressure, cardiovascular events, mortality and renal parameters

**Purpose**

Treatment

**Inclusion/Exclusion criteria****Inclusion criteria:**

Patients undergoing coronary artery bypass surgery Not being allergic to empagliflozin

**Exclusion criteria:**

Patients who cannot get enough information with echocardiography History of taking pioglitazone at least 8 weeks before entering the study Glomerular filtration rate less than 30 mg per minute per 1.73 square meters of body surface based on the CKD-EPI formula Pregnant or lactating women

**Age**

No age limit

**Gender**

Both

**Phase**

3

**Groups that have been masked**

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

**Sample size**

Target sample size: 50

**Randomization (investigator's opinion)**

Randomized

**Randomization description**

The patients are divided into 25-person groups.using the statistical software,25 out of 50 patients are randomly assigned to treatment group and 25 to placebo group.

Unit of randomization: Individual. Tools used in randomization:computer software

**Blinding (investigator's opinion)**

Double blinded

**Blinding description**

In this study, the participants, the principle investigator, the physicians and the data collectors are blinded. The patients who are going to receive the drug or placebo , are determined by numbers and these numbers are given to the head nurse and nurses of surgery department.

**Placebo**

Used

**Assignment**

Parallel

**Other design features****Secondary Ids**

empty

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

Ethics Committee of Yazd University of Medical Sciences

**Street address**

Faculty of Pharmacy,Yazd, Shohada gomnam St., Alam Square

**City**

Yazd

**Province**

Yazd

**Postal code**

8915173149

**Approval date**

2023-06-26, 1402/04/05

**Ethics committee reference number**

IR.SSU.MEDICINE.REC.1402.139

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

Patients undergoing coronary artery bypass surgery

**ICD-10 code**

T82.9

**ICD-10 code description**

Unspecified complication of cardiac and vascular prosthetic device, implant and graft

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

Reduction of blood pressure

**Timepoint**

Daily

**Method of measurement**

Mercury pressure gauge

**Secondary outcomes**

empty

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

Intervention group:Recipients of empagliflozin 10 mg

tablets from Abidi Pharmaceutical Company receive one empagliflozin tablet orally a few hours after coronary artery bypass surgery to reduce blood pressure, cardiovascular events, mortality, and control kidney parameters

**Category**

Treatment - Drugs

**2****Description**

Control group: The recipients of a placebo prepared in the pharmaceutical laboratory of the Yazd Faculty of Pharmacy in the same size and color as the empagliflozin tablet, a few hours after coronary bypass surgery, take a 10 mg tablet orally to reduce blood pressure, cardiovascular events, mortality and Renal controls

**Category**

Placebo

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

Afshar Hospital

**Full name of responsible person**

Ehsan Mirzaei

**Street address**

Yazd-Jomhuri Blvd

**City**

Yazd

**Province**

Yazd

**Postal code**

۸۹۱۷۹۴۵۵۵۶

**Phone**

+98 35 3525 5011

**Email**

ehsan.mirzaei.1369@gmail.com

**Sponsors / Funding sources****1****Sponsor****Name of organization / entity**

Yazd University of Medical Sciences

**Full name of responsible person**

Parisa Naseri

**Street address**

Faculty of pharmacy, Yazd, Shohada gomnam St.,  
Alam Square

**City**

Yazd

**Province**

Yazd

**Postal code**

8915173149

**Phone**

+98 936 850 6994

**Email**

Parisa2473@yahoo.com

**Grant name****Grant code / Reference number****Is the source of funding the same sponsor organization/entity?**

Yes

**Title of funding source**

Yazd University of Medical Sciences

**Proportion provided by this source**

1

**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**

Yazd University of Medical Sciences

**Full name of responsible person**

Ehsan Mirzaei

**Position**

Assistant Professor

**Latest degree**

Ph.D.

**Other areas of specialty/work**

Medical Pharmacy

**Street address**

Yazd Faculty of Pharmacy, Shohada Gomnam St.,  
Alam Square

**City**

Yazd

**Province**

Yazd

**Postal code**

8915173149

**Phone**

+98 35 3820 3419

**Email**

ehsan.mirzaei.1369@gmail.com

**Person responsible for scientific inquiries****Contact****Name of organization / entity**

Yazd University of Medical Sciences

**Full name of responsible person**

Ehsan Mirzaei

**Position**

Assistant Professor

**Latest degree**

Ph.D.

**Other areas of specialty/work**

Medical Pharmacy

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Yazd

**Postal code**

8915173149

**Phone**

+98 35 3820 3419

**Email**

ehsan.mirzaei.1369@gmail.com

**Person responsible for updating data****Contact****Name of organization / entity**

Yazd University of Medical Sciences

**Full name of responsible person**

Parisa Naseri

**Position**

Student

**Latest degree**

Medical doctor

**Other areas of specialty/work**

Medical Pharmacy

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8915173149

**Phone**

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**Email**

ehsan.mirzaei.1369@gmail.com

**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