

# Clinical Trial Protocol

## Iranian Registry of Clinical Trials

11 Jul 2026

**A comparison of the effectiveness, side effects and acceptability of Plavix and Osivix as anti platelet tablets in patients undergoing Coronary artery by pass.**

### Protocol summary

#### Summary

Anticoagulant effect of clopidogrel is utmost importance in coronary artery disease, especially in prevention of coronary stent thrombosis. Recently, a new local brand of clopidogrel has been launched, as Osvix® (by OSVEH Company, Tehran, Iran). This research is conducted by the aim in order to compare two locally prepared clopidogrel brands (Osvix® & Plavix®), in terms of the effect on inhibition of platelet aggregation in patients with coronary artery disease. This is a double blind randomized study. Sample population consisting of 80 patients, is admitted at Ekbatan Hospital (Hamadan, Iran) for the management of coronary artery disease. Platelet aggregation tests of all these patients will measure (by Iran Blood transfusion organization) by factors of PRP, ADP and placket count. Patients received Plavix® and Osvix® treatments regimens for one month periodically after by-pass coronary surgery.

### General information

#### Acronym

#### IRCT registration information

IRCT registration number: **IRCT201112178428N1**  
Registration date: **2014-04-19, 1393/01/30**  
Registration timing: **retrospective**

Last update:

Update count: **0**

#### Registration date

2014-04-19, 1393/01/30

#### Registrant information

##### Name

Behzad Imani

##### Name of organization / entity

Science Hamadan University of Medical

#### Country

Iran (Islamic Republic of)

#### Phone

+98 81 1838 1017

#### Email address

b.imani@umsha.ac.ir

#### Recruitment status

**Recruitment complete**

#### Funding source

Hamadan University of Medical Sciences

#### Expected recruitment start date

2012-02-05, 1390/11/16

#### Expected recruitment end date

2013-08-04, 1392/05/13

#### Actual recruitment start date

empty

#### Actual recruitment end date

empty

#### Trial completion date

empty

#### Scientific title

A comparison of the effectiveness, side effects and acceptability of Plavix and Osivix as anti platelet tablets in patients undergoing Coronary artery by pass.

#### Public title

A comparison of the efficacy of Plavix and Osivix in patients undergoing Coronary artery by pass

#### Purpose

Treatment

#### Inclusion/Exclusion criteria

Inclusion criteria: All patients undergoing open heart surgery  
Exclusion criteria: 1- Patients with a prior events of acute coronary syndrome, 2- Hepatic insufficiency, 3- History of significant bleeding disorder, 4- Those already taking anti-platelet and/or anticoagulant therapy, 5- Age less than 20 and over than 75.

**Age**

From **20 years** old to **75 years** old

**Gender**

Both

**Phase**

3

**Groups that have been masked**

*No information*

**Sample size**

Target sample size: **80**

**Randomization (investigator's opinion)**

Randomized

**Randomization description****Blinding (investigator's opinion)**

Double blinded

**Blinding description****Placebo**

Not used

**Assignment**

Parallel

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

empty

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

Hamadan University of Medical Sciences

**Street address**

Reseach Deputy, Hamadan University of Medical  
Scinces, Hamadan, Iran.

**City**

Hamadan

**Postal code**

6518133265

**Approval date**

2012-12-17, 1391/09/27

**Ethics committee reference number**

3214 /9 /35 /16

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

Heart disease

**ICD-10 code**

I97.1

**ICD-10 code description**

Other functional disturbances following cardiac surgery

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

Platelet-rich plasma

**Timepoint**

30 days after heart surgery

**Method of measurement**

Aggregometry

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

Adenosine diphosphate

**Timepoint**

30 days after heart surgery

**Method of measurement**

Aggrigometry

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

at this clinical trial study, the control group consists of 40 patients recieve Plavix tablets as an antiplatelet agent. The prescribed dose is 75 mg of Clopidogrel per day for each group. It should be noted that all patients receive 100 mg aspirin tablet daily. In order to implementation of Ex-vivo analysis, blood is mixed with 3.8% citrate and platelet aggregometry is conducted. The amount of Adenosine diphosphate, platelet-rich plasma and platelet count is measured as indicating factors of aggrigation.

**Category**

Treatment - Drugs

**2****Description**

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**Category**

Treatment - Drugs

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

Ekbatan Hospital, Hamadan,Iran.

**Full name of responsible person**

Behzad Imani

**Street address**

Operating-room Departemnt, Hamadan University of  
Medical Sciences, Mahdiah St., Hamadan, Iran.

**City**  
Hamadan

## Sponsors / Funding sources

1

### Sponsor

**Name of organization / entity**  
Research Deputy of Hamadan University of Medical Sciences

**Full name of responsible person**  
Behzad Imani

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Operating-room Departemnt, Hamadan University of Medical Sciences, Mahdieh St., Hamadan, Iran.

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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**  
Research Deputy of Hamadan University of Medical Sciences

**Proportion provided by this source**  
100

**Public or private sector**  
*empty*

**Domestic or foreign origin**  
*empty*

**Category of foreign source of funding**  
*empty*

**Country of origin**

**Type of organization providing the funding**  
*empty*

## Person responsible for general inquiries

### Contact

**Name of organization / entity**  
Hamadan University of Medical Sciences

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Behzad Imani

**Position**  
Faculty member, Master of Sciences in Nursing

**Other areas of specialty/work**

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## Person responsible for scientific inquiries

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## Person responsible for updating data

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**Other areas of specialty/work**

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**Web page address**

## Sharing plan

**Deidentified Individual Participant Data Set (IPD)**  
*empty*

**Study Protocol**  
*empty*

**Statistical Analysis Plan**  
*empty*

**Informed Consent Form**  
*empty*

**Clinical Study Report**  
*empty*

**Analytic Code**  
*empty*

**Data Dictionary**

*empty*