

Clinical Trial Protocol

Iranian Registry of Clinical Trials

02 Jun 2026

The comparison of two methods of sandbag and air bag (femosup) in reduction of coronary artery angioplasty complications

Protocol summary

Summary

The purpose of this study is comparison of two methods of sandbag and air bag in reduction of coronary artery angioplasty complications after removal of sheath. Inclusion criteria are having sufficient consciousness to work with staff and have arterial sheath in the right leg. Exclusion criteria are opiate addiction, psychological disease, existence of low back pain, blood and clotting diseases. This quasi experimental study will be done on 72 patients undergoing angioplasty who hospitalized in intensive care unit. Patients will allocate to two groups of airbag and sand bag. In the sand bag group after pulling out the sheath, routine care takes place and sand bag is put on the place of artery's puncture for 6 hours. In the airbag group after the removal of the sheath the designed device (air bag) will be fastened. The extent of hematoma, the amount of tiredness, low back pain, sedative and analgesic intake and sleep status will be evaluated in two groups

General information

Acronym

IRCT registration information

IRCT registration number: **IRCT201206306342N2**
Registration date: **2012-10-08, 1391/07/17**
Registration timing: **registered_while_recruiting**

Last update:

Update count: **0**

Registration date

2012-10-08, 1391/07/17

Registrant information

Name

Ali Asghar Ghods

Name of organization / entity

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Recruitment status

Recruitment complete

Funding source

Semnan University of Medical Sciences

Expected recruitment start date

2012-06-21, 1391/04/01

Expected recruitment end date

2012-10-21, 1391/07/30

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

The comparison of two methods of sandbag and air bag (femosup) in reduction of coronary artery angioplasty complications

Public title

effect of air bag (femosup) in reducing complication after angioplasty

Purpose

Supportive

Inclusion/Exclusion criteria

Inclusion criteria: Having sufficient consciousness to work with staff; have Arterial sheath in the right leg; use sheath no 6; received heparin as anticoagulant; arterial sheath is removed 5-6 hours after injection of heparin bolus dose. Exclusion criteria: Opiate addiction; psychological disease; previous use of analgesics and sedatives or tranquilizers during angioplasty; Integrilin received during angioplasty; previous used of Varfarin;

angioplasty due to myocardial infarction; hematoma forms before extraction of the sheath; existence of musculoskeletal disease and low back pain; blood and hepatic diseases; peripheral vascular disease; Clotting problems; existence of venous sheath; To access the femoral artery needle is made more than once; systolic blood pressure is above 180 mm Hg; diastolic BP is above 100 mm Hg; use of Hemcan glue after extraction of the sheath.

Age

No age limit

Gender

Both

Phase

N/A

Groups that have been masked

No information

Sample size

Target sample size: **72**

Randomization (investigator's opinion)

Randomized

Randomization description

Blinding (investigator's opinion)

Not blinded

Blinding description

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Semnan University of Medical Sciences

Street address

School of Nursing and Allied Health, Semnan University of Medical Sciences, Semnan, Iran

City

Semnan

Postal code

Approval date

2012-05-27, 1391/03/07

Ethics committee reference number

2744/16/91

Health conditions studied

1

Description of health condition studied

Atherosclerotic heart disease

ICD-10 code

I25.1

ICD-10 code description

Atherosclerotic heart disease

Primary outcomes

1

Description

low back pain

Timepoint

begining of intervention, 3 and 6 hours after intervention

Method of measurement

numeric rating scale of pain

2

Description

fatigue

Timepoint

begining of intervention, 3 and 6 hours after intervention

Method of measurement

numeric rating scale of fatigue

3

Description

Groin discomfort

Timepoint

begining of intervention, 3 and 6 hours after intervention

Method of measurement

numeric rating scale of groin discomfort

4

Description

hematoma

Timepoint

hours of 1,2,3,4,5,6

Method of measurement

Christensen scale

5

Description

sedative intake

Timepoint

18 hours after angioplasty

Method of measurement

Patient records

6

Description

opiate intake

Timepoint

18 hours after angioplasty

Method of measurement

patient records

7

Description

sleep status

Timepoint

Day after angioplasty

Method of measurement

questionnaire

Secondary outcomes

empty

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

use of air bag (femosup) immediatly after removal of sheath

Category

Other

2**Description**

use of sand bag immediatly after removal of sand bag

Category

Other

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

Razavi Hospital

Full name of responsible person

Samira Bagheri

Street address

Razavi Hospital, Mashhad

City

Mashhad

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

Semnan University of Medical Sciences

Full name of responsible person

Raheb Ghorbani

Street address

Vice Chancellor for Research, Semnan University of Medical Sciences, Semnan

City

Semnan

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

Yes

Title of funding source

Semnan 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**

Semnan University of Medical Sciences

Full name of responsible person

Ali Asghar Ghods

Position

faculty member

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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