

Clinical Trial Protocol

Iranian Registry of Clinical Trials

20 Jun 2026

Efficacy of Silver based antiseptics application for control of Central Venous catheters colonization in cardiac surgery

Protocol summary

Summary

Central venous (CV) catheters play an essential role in the management of critically ill patients in the Intensive Care Unit (ICU). CV lines are, however, allied to catheter-associated blood stream infections. Bacterial colonization of CV lines is deemed the main cause of catheter-associated infection. The purpose of our study is to compare bacterial colony counts in the catheter site before CV line insertion in two groups of post-cardiac surgery patients: a group receiving Sanosil (an antiseptic agent composed of H₂O₂ and silver) and a control group. The study is design as an interventional prospective double-blinded clinical trial recruited the patients in three post-cardiac surgery ICU's of a heart center. 250 patients will be randomized to be treated with sanosil plus their routine infection control prophylactic measures or without it. The participants will divided into interventional and control groups. Sanosil spray will be added to the routine preparation procedure (Chlorhexidine bath one day before and scrub with Povidone-Iodine just before the CV line insertion). After the removal of the CV lines at an average of three days, the catheters tips will be sent for culture and evaluation of colony counts.

General information

Acronym

IRCT registration information

IRCT registration number: **IRCT201009014672N1**
Registration date: **2013-05-25, 1392/03/04**
Registration timing: **retrospective**

Last update:

Update count: **0**

Registration date

2013-05-25, 1392/03/04

Registrant information

Name

Fardin Yousefshahi

Name of organization / entity

Tehran University of Medical Sciences

Country

Iran (Islamic Republic of)

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+98 21 8802 9600

Email address

yousefshahi@tums.ac.ir

Recruitment status

Recruitment complete

Funding source

Tehran University of Medical Sciences

Expected recruitment start date

2013-03-21, 1392/01/01

Expected recruitment end date

2013-05-22, 1392/03/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Efficacy of Silver based antiseptics application for control of Central Venous catheters colonization in cardiac surgery

Public title

Efficacy of Silver based antiseptics application for decrease of Central Venous catheter infections

Purpose

Prevention

Inclusion/Exclusion criteria

Inclusion criteria: candidate for open-heart surgery in Tehran Heart Center; signing the consent for participation in the study; without any evidence of

infection at the time of admission. Exclusion criteria: infection on arrival or occurrence of infection in first 48 hours after arrival; not being admitted to the post ICU for any reason.

Age

No age limit

Gender

Both

Phase

N/A

Groups that have been masked

No information

Sample size

Target sample size: **250**

Randomization (investigator's opinion)

Randomized

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

Double blinded

Blinding description**Placebo**

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Tehran University of Medical Sciences

Street address

PO Box 1417614411, Floor 8, Tehran University of Medical Sciences, Qods Street, Tehran, Iran.

City

Tehran

Postal code**Approval date**

2010-12-12, 1389/09/21

Ethics committee reference number

130/1122/89/3

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

catheter related infection profilaxis

ICD-10 code

B99

ICD-10 code description

Other and unspecified infectious diseases

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

colonization of bacteria in Central Venous Catheter

Timepoint

after catheter removal

Method of measurement

positive or negative cultures

2**Description**

the type of isolated microorganism from the CVline

Timepoint

after catheter removal

Method of measurement

the type of microorganism

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

leuckocytosis

Timepoint

admission period

Method of measurement

wight blood cell count more than normal value

2**Description**

Fever

Timepoint

admission period

Method of measurement

body temperature more than 37.8 centigrade

3**Description**

sepsis symptoms and signs

Timepoint

admission period

Method of measurement

evaluation of clinical and para-clinical factores

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

Sanosil spraying on the insertion point by an standard spray, 3 puffs, half an hour before surgery, in preparation room

Category

Treatment - Drugs

2

Description

distilled water spraying on the insertion point by an standard spray, 3 puffs, half an hour before surgery, in preparation room

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Tehran Heart Center

Full name of responsible person

Fardin Yousefshahi

Street address

Tehran Heart Center, Al-ahmad & Kargar Cross

City

Tehran

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Tehran Heart Center, Infection Control Section

Full name of responsible person

Dr. Mohamadali Broomand

Street address

Infection Control section, Tehran Heart Center, Al-ahmad & Kargar Cross, Tehran, Iran.

City

Tehran

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Tehran Heart Center, Infection Control Section

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

Tehran University of Medical Sciences

Full name of responsible person

Fardin Yousefshahi MD

Position

Assistant Professor

Other areas of specialty/work

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Person responsible for updating data

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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