

Clinical Trial Protocol

Iranian Registry of Clinical Trials

11 Jul 2026

comparison of success rate and complications of inguinal and subinguinal varicocelelectomy operation

Protocol summary

Summary

In this clinical trial 64 patients older than 18 years old suffering from varicocele referred to urology clinic candidate for surgical varicocelelectomy randomly allocated in two groups. In both groups after taking informed consent form and approval of ethics committee, diagnosis of varicocele will be confirmed by physical examination or ultrasonography and semen analysis will be done for all of them. In presence of infertility, pain or abnormal sperm analysis varicocelelectomy will be performed by surgical method without using loops or microscope. In first group (subinguinal) under general anesthesia in supine position with a 3 centimeter transverse incision on external inguinal ring skin and subcutaneous fat dissected and after separating vas deference and lymphatics, spermatic veins dissect and ligate by 3-0 silk suture. In second group skin will be incised as first group, but with incision on external oblique fascia cord mobilized and operation will done as first group. In both groups patients discharge after operation and will be followed one week next and exam for presence of hematoma, wound infection and pain by visual analogue scale. One and 3 months later they visit again and check by history, physical examination or ultrasonography for presence of hydrocele and recurrence of varicocele. After three month spermogram will be checked in all of them for success of operation and improvement of infertility. All data will be recorded and analyzed by SPSS software version 21 and analytical descriptive and Qi square tests.

General information

Acronym

IRCT registration information

IRCT registration number: **IRCT201710131323N12**

Registration date: **2017-10-20, 1396/07/28**

Registration timing: **registered_while_recruiting**

Last update:

Update count: **0**

Registration date

2017-10-20, 1396/07/28

Registrant information

Name

Sadrollah Mehrabi

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 74 3334 6070

Email address

dr.mehrabi@yums.ac.ir

Recruitment status

Recruitment complete

Funding source

Research vice chancellor of Yasuj University of Medical Sciences

Expected recruitment start date

2016-10-21, 1395/07/30

Expected recruitment end date

2017-10-22, 1396/07/30

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

comparison of success rate and complications of inguinal and subinguinal varicocelelectomy operation

Public title

comparison of inguinal and subinguinal varicocelelectomy

Purpose

Treatment

Inclusion/Exclusion criteria

inclusion criteria:males suffering from varicocele with pain and impairment of spermogram aged more than 18
Exclusion criteria:noncooperation for followup,presence of morbid obesity with BMI >35,coincidence of inguinal hernia with varicocele,history of previous varicocelectomy and recurrence

Age

From **18 years** old to **60 years** old

Gender

Male

Phase

N/A

Groups that have been masked

No information

Sample size

Target sample size: **64**

Randomization (investigator's opinion)

N/A

Randomization description

Blinding (investigator's opinion)

Single blinded

Blinding description

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

1

Registry name

-

Secondary trial Id

-

Registration date

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics committee of Yasuj University of Medical Sciences

Street address

Yasuj University of Medical Sciences, Mottahari srteet, Yasuj, Iran

City

Yasuj

Postal code

Approval date

2016-12-30, 1395/10/10

Ethics committee reference number

IR.YUMS.REC.2016.150

Health conditions studied

1

Description of health condition studied

varicocele

ICD-10 code

186-1

ICD-10 code description

scrotal varices

Primary outcomes

1

Description

improvement of fertility and spermogram

Timepoint

befor and 3 months after operation

Method of measurement

with semen analysis

Secondary outcomes

1

Description

Pain relief

Timepoint

one week and one month after operation

Method of measurement

visit and examinatin and check by visual analogue scale

Intervention groups

1

Description

performing subinguinal varicocelectomy in supine position without using optic loop

Category

Treatment - Drugs

2

Description

performing inguinal varicocelectomy in supine position without using optic loop

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Shaheed Beheshti hospital

Full name of responsible person

Sadrollah Mehrabi

Street address

Shaheed Beheshi hospital, Montazeri street, Yasuj,Iran

City

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Research vice chancellor of Yasuj University of Medical Sciences

Full name of responsible person

Dr Ali Mousavizadeh

Street address

Mottahari street, Research vice chancellor, Yasuj University of Medical Sciences, , Iran

City

Yasuj

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Research vice chancellor of Yasuj 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

Faculty of Medicine, Yasuj University of Medical Sciences

Full name of responsible person

Sadrollah Mehrabi

Position

professor, Fellowship of endourology and laparoscopy

Other areas of specialty/work

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Person responsible for scientific inquiries

Contact

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Full name of responsible person

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Other areas of specialty/work

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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

