

Clinical Trial Protocol

Iranian Registry of Clinical Trials

09 Jun 2026

Evaluation of the efficacy of dilapane-S gel in comparison with misoprostol during pre-hysteroscopic cervical preparation

Protocol summary

Study aim

Evaluation of the efficacy of Dilapan S gel in comparison with misoprostol for cervical preparation before hysteroscopy

Design

It is a double-blind clinical trial performed on 120 patients undergoing hysteroscopy. Randomization is performed simply and patients are divided into control and control groups with parallel groups

Settings and conduct

This study will be performed in Hazrat Rasool Akram Hospital based on the prospective method. The study population is female patients referred to the gynecology clinic in the mentioned hospital who are candidates for hysteroscopic surgery. According to the sample size calculations, 120 patients with the mentioned conditions were selected and after providing the necessary explanations about this The plan will receive written consent.

Participants/Inclusion and exclusion criteria

Inclusion criteria included postmenopausal women (at least more than 1 year after the last menstrual cycle) or Noliplar or Multipar who have no history of normal delivery who are candidates for hysteroscopic surgery due to uterine myoma of uterine polyp, increased endometrial thickness confirmed by ultrasound or Abnormal vaginal bleeding. Exclusion criteria are contraindications to hysteroscopy, recent or current pelvic infection, previous surgery on the cervix, confirmed cervical or breast malignancies, history of vaginal delivery, sensitivity to misoprostol and prostaglandins.

Intervention groups

Patients in the intervention group underwent dilapane S (3MM * 55MM) 6 hours before the operation and vaginal misoprostol was placed 6 hours before the operation in the control group

Main outcome variables

Dilatation, pain intensity and bleeding of the patient

after hysteroscopy

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20191123045476N2**

Registration date: **2021-02-19, 1399/12/01**

Registration timing: **registered_while_recruiting**

Last update: **2021-02-19, 1399/12/01**

Update count: **0**

Registration date

2021-02-19, 1399/12/01

Registrant information

Name

Samaneh Rokhgireh

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 21 6650 9283

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2020-09-22, 1399/07/01

Expected recruitment end date

2021-03-19, 1399/12/29

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Evaluation of the efficacy of dilapane-S gel in comparison with misoprostol during pre-hysteroscopic cervical preparation

Public title

Evaluation of the effect of Dilapane-S gel on preoperative cervical preparation

Purpose

Supportive

Inclusion/Exclusion criteria

Inclusion criteria:

Postmenopausal women (at least 1 year after the last menstrual cycle) or NoliPar or MultiPar who have no history of natural childbirth are candidates for hysteroscopic surgery due to uterine myoma and uterine polyp. Postmenopausal women (at least more than 1 year after the last menstrual cycle) or Nolie Par or MultiPar who have no history of normal delivery are candidates for hysteroscopic surgery due to increased endometrial thickness confirmed by ultrasound or abnormal vaginal bleeding

Exclusion criteria:

Contraindications to hysteroscopy Recent or current pelvic infections Previous cervical surgery Confirmed cervical or breast malignancies Vaginal delivery history Hypersensitivity to misoprostol and prostaglandins

Age

From **49 years** old to **52 years** old

Gender

Female

Phase

3

Groups that have been masked

- Outcome assessor
- Data analyser

Sample size

Target sample size: **120**

Randomization (investigator's opinion)

Randomized

Randomization description

In this study, restricted randomization sampling method is used that the study groups have equal sample size. Random allocation rule is one of the limited random sampling methods used in the present study. And patients are equally divided into two groups. On how to randomize, researchers first determine a total sample size, then randomly assign a set of them to group A and the rest to group B. For example, in a study with a sample size of 200 people, 100 balls For intervention group A and 100 balls, for intervention group B, it is placed in a lottery container and then the balls are randomly removed from the container without replacement and the created sequence is recorded. This method is used for two or more group trials.

Blinding (investigator's opinion)

Double blinded

Blinding description

Patients in the intervention group are placed with

dilapane 6 hours before hysteroscopy. 6 hours later, before hysteroscopy, dilapane is removed and the patient is transferred to the operating room. In the control group, 6 hours before surgery (the night before surgery) will receive 200 micrograms of vaginal misoprostol tablets (Sami Saz Pharmaceutical Company). How medications are prescribed to the surgeon who evaluates the outcome will be kept secret. The analyzer is also unaware of the type of medication being administered.

Placebo

Not used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethics Committee of Iran University of Medical Sciences

Street address

Sattar Khan St., Maziar Mansouri St., Rasoul Akram Hospital

City

Tehran

Province

Tehran

Postal code

1445613131

Approval date

2020-11-07, 1399/08/17

Ethics committee reference number

IR.IUMS.REC.1399.764

2

Ethics committee

Name of ethics committee

Ethics Committee of Iran University of Medical Sciences

Street address

Sattar Khan St., Maziar Mansouri St., Rasoul Akram Hospital

City

Tehran

Province

Tehran

Postal code

1445613131

Approval date

2020-11-07, 1399/08/17

Ethics committee reference number

IR.IUMS.REC.1399.764

Health conditions studied

1

Description of health condition studied

Hysteroscopy

ICD-10 code

ICD-10 code description

Primary outcomes

1

Description

Cervical dilatation

Timepoint

The time required to start dilatation until the arrival of a hysteroscope or resectoscope will be recorded as a conclusion

Method of measurement

Initially, the initial condition of the cervix, including the position and softness of the cervix, will be evaluated observably. No resistance is passed as dilatation before hysteroscopy and will be recorded as a conclusion. Our ultimate goal will be dilatation with Hagar number 10-8.

Secondary outcomes

1

Description

Severe pain and bleeding after hysteroscopy

Timepoint

Immediately after the patient regains consciousness

Method of measurement

visual analog scale, which includes a table to evaluate the patient's pain and bleeding in the form of zero indicates painless and ten indicates severe and unbearable pain and bleeding.

Intervention groups

1

Description

Intervention group: Patients in the intervention group undergoing placement with Dilapane S (3MM * 55MM) made by Kardan Elixir Company, 6 hours before hysteroscopy, which is first moistened with sterile water or saline, and then take the Dilapane S handle gently and without Excess pressure is applied to the cervical canal only when it passes through the external and internal holes. The Dilapan handle should be placed around the external hole and not go beyond it, and 6 hours later and before hysteroscopy to remove the Dilapan S handle only with Forceps are taken and a continuous downward pull is applied along the longitudinal axis of the dilator, which makes it easy and painless to exit.

Category

Treatment - Devices

2

Description

Control group: In the control group, 6 hours before surgery (the night before surgery) will receive 200 micrograms of vaginal misoprostol tablets (Sami Saz Pharmaceutical Company).

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Rasool Akram Hospital

Full name of responsible person

Samaneh Rokhgireh

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Starkhan.Mansouri Street. Rasoul Akram Hospital

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Iran University of Medical Sciences

Full name of responsible person

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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

Iran University of Medical Sciences

Proportion provided by this source

100

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**

Iran University of Medical Sciences

Full name of responsible person

Samaneh Rokhgireh

Position

Associate professor

Latest degree

Specialist

Other areas of specialty/work

Gynecology and Obstetrics

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Samaneh Rokhgireh

Position

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Latest degree

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Other areas of specialty/work

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

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Sharing plan**Deidentified Individual Participant Data Set (IPD)**

Yes - There is a plan to make this available

Study Protocol

Yes - There is a plan to make this available

Statistical Analysis Plan

Yes - There is a plan to make this available

Informed Consent Form

Yes - There is a plan to make this available

Clinical Study Report

Yes - There is a plan to make this available

Analytic Code

Yes - There is a plan to make this available

Data Dictionary

Yes - There is a plan to make this available

Title and more details about the data/document

Following the publication of the article, confidential information such as patient profile and Hospital, ... deletion and other information Will be made available to researchers

When the data will become available and for how long

After publishing the article

To whom data/document is available

Medical specialists

Under which criteria data/document could be used

Medical professionals can turn to data for research purposes gain access

From where data/document is obtainable

Refer to the email of the responsible author

What processes are involved for a request to access data/document

Official and academic email to the responsible author

Comments