

Clinical Trial Protocol

Iranian Registry of Clinical Trials

27 Aug 2026

The comparative evaluation of the effect of sublingual and cervical misoprostol on the duration of abortion in patients candidate for abortion in the first trimester of pregnancy

Protocol summary

Study aim

The Determination of the effect of sublingual and cervical misoprostol on the the abortion rate and duration of abortion in the first trimester of pregnancy in patients candidate for abortion

Design

Clinical trials , with parallel groups, randomized

Settings and conduct

This study is performed in Qazvin University of Medical Sciences. In this study, all pregnant women with a gestational age of 1 to 12 weeks who are candidates for legal abortion and referred to Kowsar and Mehregan hospitals in Qazvin are examined and included in the study. This study is performed as a randomized single-blind clinical trial (participant in the project). Data collection is completed by the questioner which is done through a checklist. The sample size was 200 people. If they meet the inclusion criteria, enter the study and are randomly divided into two equal groups a and b.

Participants/Inclusion and exclusion criteria

Inclusion Criteria : Unifetal pregnancy , Missed abortion , Fetal malformations , Blighted ovum, Trisomies , Gestational age (12 weeks and less than 12 weeks) ; Exclusion Criteria : Rupture of membrane , History of cesarean section more than once, Multifetal pregnancy, History of allergy to misoprostol and prostaglandins , Contraindications of prostaglandin use (like heart disease, kidney disease, history of seizures and hypotension)

Intervention groups

At the beginning , patients of group A will receive 400 micrograms of sublingual misoprostol along with cervical placebo and , patients of group b will receive 400 micrograms of wet ted cervical misoprostol along with sublingual placebo, and then the dose is repeated every 4 hours until the abortion, and the maximum duration of administration of the drug will be up to 48 hours.

Main outcome variables

The time interval between induction of abortion and excretion of pregnancy products

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20200127046280N1**

Registration date: **2020-10-16, 1399/07/25**

Registration timing: **prospective**

Last update: **2020-10-16, 1399/07/25**

Update count: **0**

Registration date

2020-10-16, 1399/07/25

Registrant information

Name

Amirhossein Gholamlou

Name of organization / entity

Country

Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2020-10-22, 1399/08/01

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

The comparative evaluation of the effect of sublingual and cervical misoprostol on the duration of abortion in patients candidate for abortion in the first trimester of pregnancy

Public title

The comparative evaluation of the effect of sublingual and cervical misoprostol on first trimester abortion

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Unifetal pregnancy Missed abortion Fetal malformations blighted ovum Trisomies Gestational age (12 weeks and less than 12 weeks)

Exclusion criteria:

Rupture of membrane History of cesarean section more than once Multifetal pregnancy History of allergy to misoprostol and prostaglandins Contraindications of prostaglandin use (like heart disease, kidney disease, history of seizures and hypotension)

AgeFrom **16 years** old to **45 years** old**Gender**

Female

Phase

2-3

Groups that have been masked

- Participant

Sample sizeTarget sample size: **200****Randomization (investigator's opinion)**

Randomized

Randomization description

Patients who meet the inclusion criteria are divided into two intervention groups using simple randomization method. The randomization method used in this study is the use of a table of random numbers. Random number table is a set of numbers that is generated without a specific pattern or order and they generated randomly and they are formed in a table. In first the direction of reading the numbers was specified. To read the numbers, random numbers are read from the left side of the table, then even numbers extracted from the table are allocated to intervention group a and odd numbers extracted from the table are allocated to intervention group b.

Blinding (investigator's opinion)

Single blinded

Blinding description

Participants are unaware of treatment allocation and are divided into groups a and b. Group a receives sublingual misoprostol with cervical placebo and group b receives cervical misoprostol with sublingual placebo. Finally, the results of the work are filled by the midwife of the ward,

using the questionnaire form and provided to the researcher.

Placebo

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Ethics committee of Qazvin University Of Medical Science

Street address

No 4, mavedat ave, shahid beheshti blvd

City

Qazvin

Province

Qazvin

Postal code

1391134156

Approval date

2020-01-25, 1398/11/05

Ethics committee reference number

IR.QUMS.REC.1398.280

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

Abortion

ICD-10 code

O02.1

ICD-10 code description

Missed abortion

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

The time interval between induction of abortion and excretion of pregnancy products

Timepoint

After intervention and then every one hour

Method of measurement

clock

2**Description**

Nausea (drug side effects)

Timepoint

After intervention and then every one hour

Method of measurement

Clinical examination and questioning of the patient

3**Description**

Vomiting (drug side effects)

Timepoint

After intervention and then every one hour

Method of measurement

Clinical examination and questioning of the patient

4**Description**

Fever (drug side effects)

Timepoint

After intervention and then every one hour

Method of measurement

Thermometer

5**Description**

Diarrhea (drug side effects)

Timepoint

After intervention and then every one hour

Method of measurement

Clinical examination and questioning of the patient

6**Description**

Headache (drug side effects)

Timepoint

After intervention and then every one hour

Method of measurement

Clinical examination and questioning of the patient

7**Description**

abortion type (Complete or incomplete)

Timepoint

After intervention and then every one hour

Method of measurement

Clinical examination and questioning of the patient

Secondary outcomes

empty

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

First intervention group (group a): This group will receive 400 mcg of sublingual misoprostol from the brand (Cytotec, Searle, England) along with cervical placebo and then the mentioned dose will be repeated every 4 hours until the abortion. The maximum duration of drug administration will be up to 48 hours.

Category

Treatment - Drugs

2**Description**

Intervention group 2 (group b): This group will receive 400 mcg of cervical misoprostol from brand (Cytotec, Searle, England) moistened with a few drops of distilled water along with sublingual placebo and then the mentioned dose is repeated every 4 hours until abortion. The maximum duration of drug administration will be up to 48 hours.

Category

Treatment - Drugs

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

Kowsar hospital

Full name of responsible person

Masoumeh Dadashaliha

Street address

Kowsar Hospital, Kowsar Alley, next to the electricity office, Taleghani St

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2**Recruitment center****Name of recruitment center**

Mehregan hospital

Full name of responsible person

Masoumeh Dadashaliha

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Mehregan Hospital, next to Dialysis Center, Ferdowsi St.

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Qazvin University of Medical Sciences

Full name of responsible person

Dr. Mohammad Mahdi Emamjomeh

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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

Qazvin 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

Qazvin University of Medical Sciences

Full name of responsible person

Masoumeh Dadashaliha

Position

Associate Professor

Latest degree

Specialist

Other areas of specialty/work

Gynecology and Obstetrics

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Person responsible for scientific inquiries

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Latest degree

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Other areas of specialty/work

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Person responsible for updating data

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Full name of responsible person

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Latest degree

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Other areas of specialty/work

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Sharing plan

Deidentified Individual Participant Data Set (IPD)

Yes - There is a plan to make this available

Study Protocol

Undecided - It is not yet known if there will be a plan to make this available

Statistical Analysis Plan

Not applicable

Informed Consent Form

Undecided - It is not yet known if there will be a plan to make this available

Clinical Study Report

Not applicable

Analytic Code

Not applicable

Data Dictionary

Not applicable

Title and more details about the data/document

A part of the data, such as information about the main outcome, can be shared.

When the data will become available and for how long

Access period starts 6 months after publishing the results.

To whom data/document is available

It will be available only to researchers working in academic and scientific institutions.

Under which criteria data/document could be used

If another similar clinical trial is performed

From where data/document is obtainable

Email

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

After Receiving E-mail And Proposal And Ensuring That Information Is Not Misused

Comments