

Clinical Trial Protocol

Iranian Registry of Clinical Trials

04 Jul 2026

Investigation of the effects of vaginal probiotic capsules on prolonging the time of membrane rupture up to delivery and on maternal and fetal complications.

Protocol summary

Study aim

Investigation of the effects of vaginal probiotic capsules on prolonging the time of membrane rupture up to delivery and on maternal and fetal complications

Design

In this study, 64 pregnant women who have the inclusion criteria and refer to Ghaem, Emam Reza and Ommolbanin hospitals are chosen. Participants are randomly divided to intervention and control group (Randomization is done by computer).

Settings and conduct

Regarding that the most important factor in neonatal death is Preterm delivery, and on the other hand the reason of 35 percent of these Preterm delivery is preterm membrane rupture and also due to the fact that after delivery, the more the age, the less the complications related to preterm infant, in this study. Therefore, with the prescription of vaginal probiotic capsules and maintaining normal vaginal microbial flora, it is aimed to increase pregnancy age and reduce the complications of being preterm. This study will be carried out in Ghaem, Emam Reza and Ommolbanin hospitals in Mashhad. No blinding is done and patients are randomly divided to two groups.

Participants/Inclusion and exclusion criteria

Inclusion criteria: live singleton pregnancy, being pregnant for 24 to 32 weeks, no symptoms of chorioamniotic, no sign of fetal distress, no vaginal bleeding, consent of mothers to participate in the study. Exclusion criteria: sings of chorioamniotic, disturbed fetal health assessment test, vaginal bleeding, unwillingness of patients to continue participating in the study.

Intervention groups

for randomization, two equal sets of green and blue cards are provided and placed in envelopes (Randomization is done by computer). An envelope will be allocated to each patient. In both green and blue

groups, the awaiting treatment is carried out as following: In the first 48 hours of hospitalization, 2 grams of intravenous Ampicillin injection per 6 hours and then for a week, 500-milligram Amoxicillin capsules per 8 hours and 400-milligram Erythromycin pills per 6 hours are prescribed. Also two doses of 12-milligram intramuscular Betamethasone injection within 24 hours is prescribed. Mother's vital signs and fetus health are precisely controlled through daily NST, sonography twice a week and CBC every other day. Additionally for patients who have green sets of cards, during this period of time, one vaginal probiotic capsule made by ZIST TAKHMIR company is prescribed daily for 10 days. 2 weeks after delivery, infants and mothers in both groups will be still be followed.

Main outcome variables

Main outcomes: duration of awaiting treatment and pregnancy age at the time of delivery

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20141209020264N3**

Registration date: **2018-03-19, 1396/12/28**

Registration timing: **registered_while_recruiting**

Last update: **2018-03-19, 1396/12/28**

Update count: **0**

Registration date

2018-03-19, 1396/12/28

Registrant information

Name

Seyedeh Azam Pourhoseini

Name of organization / entity

Mashhad University of Medical Science

Country

Iran (Islamic Republic of)

Phone

+98 51 3847 1446

Email address

pourhoseinia@mums.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2017-10-23, 1396/08/01

Expected recruitment end date

2019-10-23, 1398/08/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Investigation of the effects of vaginal probiotic capsules on prolonging the time of membrane rupture up to delivery and on maternal and fetal complications.

Public title

Investigation of the effects of vaginal probiotic capsules on prolonging the time of membrane rupture up to delivery and on maternal and fetal complications.

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria:

live singleton pregnancy being pregnant for 24 to 32 weeks no symptoms of chorioamniotic no sign of fetal distress no vaginal bleeding consent of mothers to participate in the study

Exclusion criteria:

sings of chorioamniotic disturbed fetal health assessment test vaginal bleeding unwillingness of patients to continue participating in the study

Age

No age limit

Gender

Female

Phase

3

Groups that have been masked

No information

Sample size

Target sample size: 64

Randomization (investigator's opinion)

Randomized

Randomization description

for randomization, two equal sets of green and blue cards are provided and placed in envelopes (Randomization is done by computer). An envelope will be allocated to each patient.

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

Ethics committee of Mashhad University of Medical Sciences

Street address

Central Building of Mashhad University of Medical Sciences (Ghorshi), Daneshgah 16, Daneshgah street

City

Mashhad

Province

Razavi Khorasan

Postal code

9176699199

Approval date

2017-09-27, 1396/07/05

Ethics committee reference number

IR.MUMS.REC.1396.194

Health conditions studied

1

Description of health condition studied

prevention of Preterm delivery and maternal and fetal complications

ICD-10 code

060.0

ICD-10 code description

Preterm labour without delivery

Primary outcomes

1

Description

prolonging the time of membrane rupture up to 34 weeks

Timepoint

Precise control of mother's vital signs and fetus health, two times of sonography during a week and CBC every other day

Method of measurement

Fetus health is evaluated through NST.

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group: for pregnant women with premature membrane rupture, in the first 48 hours of hospitalization, 2 grams of intravenous Ampicillin injection per 6 hours and then for a week, 500-milligram Amoxicillin capsules per 8 hours and 400-milligram Erythromycin pills per 6 hours are prescribed. Also two doses of 12-milligram intramuscular Betamethasone injection within 24 hours is prescribed. Mother's vital signs and fetus health are precisely controlled through daily NST, sonography twice a week and CBC every other day. Additionally during this period of time, one vaginal probiotic capsule made by ZIST TAKHMIR company is prescribed daily for 10 days.

Category

Treatment - Drugs

2

Description

Control group: for pregnant women with premature membrane rupture, in the first 48 hours of hospitalization, 2 grams of intravenous Ampicillin injection per 6 hours and then for a week, 500-milligram Amoxicillin capsules per 8 hours and 400-milligram Erythromycin pills per 6 hours are prescribed. Also two doses of 12-milligram intramuscular Betamethasone injection within 24 hours is prescribed. Mother's vital signs and fetus health are precisely controlled through daily NST, sonography twice a week and CBC every other day.

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Emam Reza hospital

Full name of responsible person

Mahnoush Rohizadeh

Street address

Emam Reza hospital, Emam Reza Sq, Ebne-Sina Ave

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9137913316

Phone

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Email

rouhizadehm941@mums.ac.ir

2

Recruitment center

Name of recruitment center

Ghaem hospital

Full name of responsible person

Seyedeh Azam Pourhoseini

Street address

Ghaem hospital, Ahmad Abad Ave

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Phone

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Email

pourhoseinia@mums.ac.ir

3

Recruitment center

Name of recruitment center

Ommol-Banin hospital

Full name of responsible person

Mahnoush Rohizadeh

Street address

Ommol-Banin hospital, Azadi 16th, Azadi Avenue

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Mashhad University of Medical Sciences

Full name of responsible person

Dr Mohsen Tafaghodi

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Central Building of Mashhad University of Medical Sciences (Ghorshi), Daneshgah 16, Daneshgah street

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Mashhad

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ramresearch@mums.ac.ir

Grant name

Grant code / Reference number

951811

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

Yes

Title of funding source

Mashhad 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**

Mashhad University of Medical Sciences

Full name of responsible person

Seyedeh Azam Pourhoseini

Position

Assistant Professor

Latest degree

Specialist

Other areas of specialty/work

Gynecology and Obstetrics

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Person responsible for scientific inquiries**Contact****Name of organization / entity**

Mashhad University of Medical Sciences

Full name of responsible person

Seyedeh Azam Pourhoseini

Position

Assistant Professor

Latest degree

Specialist

Other areas of specialty/work

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

Mashhad University of Medical Sciences

Full name of responsible person

Seyedeh Azam Pourhoseini

Position

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Latest degree

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Other areas of specialty/work

Gynecology and Obstetrics

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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

All data can be shared after patients are made unidentifiable.

When the data will become available and for how long

Data can be accessible 6 months after results are published.

To whom data/document is available

Data will be available for researchers in universities and other scientific institutes.

Under which criteria data/document could be used

Carrying out analysis on data is permitted.

From where data/document is obtainable

Data can be accessible through sending an email to the corresponding author.

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

After sending a request email to the corresponding author, data will be sent in 1 month.

Comments