

Clinical Trial Protocol

Iranian Registry of Clinical Trials

01 Sep 2026

Effects of topical magnesium sulfate on duration of labor and childbirth experience: a randomized controlled trial

Protocol summary

Study aim

To determine effect of topical magnesium sulfate on the duration of labor and childbirth experience in term pregnancy

Design

Randomized superiority placebo-controlled double-blind trial with two parallel arms on 98 women hospitalized for vaginal delivery. The randomizer software will be used for randomization.

Settings and conduct

In Taleghani teaching hospital in Tabriz-Iran, eligible women hospitalized for vaginal delivery will be randomized into one of the two study groups using stratified block randomization, after receiving informed written consent. They will be continuously monitored until delivery and followed up until one month after delivery. The drug and placebo will be identical in appearance. Participants, those recruiting and allocating participants, intervener, and data collector and analyzer will be blinded.

Participants/Inclusion and exclusion criteria

Participants will be nulliparous or women with a 1-2 parity, aged 18-35 years and body mass index 19.8-30 kg/m², without a history of cesarean section, with alive cephalic single mature fetus with estimated weight 2500 to 4000 g. Other inclusion criteria include low risk pregnancy, spontaneous or non-spontaneous onset of labor, 40-70% cervical effacement, station -2 to 0, and woman adequate literacy. Exclusion criteria include: history of infertility, risky diseases, normal delivery contraindications (including cephalo-pelvic disproportion), fetal heart rate disorders before intervention, and unwillingness to participate in the study.

Intervention groups

Intervention group: receiving 10 mL of magnesium sulfate 50% intravaginally. Control group: receiving 10 mL of distilled water intravaginally.

Main outcome variables

The interval between intervention initiation and vaginal delivery; Score of childbirth experience

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20100414003706N40**

Registration date: **2021-11-21, 1400/08/30**

Registration timing: **prospective**

Last update: **2021-11-21, 1400/08/30**

Update count: **0**

Registration date

2021-11-21, 1400/08/30

Registrant information

Name

Sakineh Mohammad-Alizadeh-Charandabi

Name of organization / entity

Country

Iran (Islamic Republic of)

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alizades@tbzmed.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2021-12-10, 1400/09/19

Expected recruitment end date

2022-05-09, 1401/02/19

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Effects of topical magnesium sulfate on duration of labor and childbirth experience: a randomized controlled trial

Public title

Comparison of effect of topical magnesium sulfate and placebo (distilled water) on duration of labor and childbirth experience

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Women aged 18-35 years, nulliparous or with 1-2 parous without a history of cesarean section Body mass index 19.8-30 kg/m² (based on weight of pre-pregnancy or first trimester of pregnancy) A live singleton term fetus (gestational age of 37-41 weeks) in estimated weight 2500 to 4000 g and cephalic presentation Having sufficient literacy to read and understand study questionnaires Spontaneous or non-spontaneous onset of labor process Cervical effacement 40-70% Station -2 to 0

Exclusion criteria:

History of infertility Contraindications to vaginal delivery including placental abruption, umbilical cord prolapse, cephalopelvic disproportion (CPD) Fetal heart rate disorders in the pre-intervention stage High risk pregnancy (bleeding in the third trimester, placental abruption, placenta previa, fetal growth disorder, etc.) Risky diseases (such as severe anemia (hemoglobin less than 7 g/dL) or blood disorders, heart disease, lung disease, connective tissue and smooth muscle problems) Unwillingness to participate in the study

Age

From **18 years** old to **35 years** old

Gender

Female

Phase

3

Groups that have been masked

- Participant
- Care provider
- Investigator
- Outcome assessor
- Data analyser

Sample size

Target sample size: **98**

Randomization (investigator's opinion)

Randomized

Randomization description

Allocation sequence will be generated using stratified block randomization (stratified by previous history of labor and type of labor onset (spontaneous or induction)) with block size of four and allocation ratio 1:1 using a computerized program (Randomizer). Sequentially numbered opaque sealed envelopes including syringe containing the drug or placebo will be used to conceal the allocation.

Blinding (investigator's opinion)

Double blinded

Blinding description

Syringes containing 10 mL magnesium sulfate or distilled water (identical in appearance) will be packed in sequentially numbered opaque sealed envelopes. The preparation of the envelopes will be performed by a person non-involved in participant recruitment and data collection. The participants, those involved in the recruitment, allocation and data collection, also analyzer will not be blinded.

Placebo

Used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Ethics committee of Tabriz University of Medical Sciences

Street address

No. 2 Central Building, Tabriz University of Medical Sciences, Golgasht Ave., Tabriz, Iran

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East Azarbaijan

Postal code

5166614766

Approval date

2021-10-25, 1400/08/03

Ethics committee reference number

IR.TBZMED.REC.1400.726

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

Prolonged labor

ICD-10 code

O63.0

ICD-10 code description

Prolonged first stage (of labor)

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

The interval between intervention initiation and childbirth

Timepoint

After delivery

Method of measurement

Timer

2**Description**

Score of maternal birth experience

Timepoint

One month after delivery

Method of measurement

Childbirth Experience Questionnaire 2.0

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

Score of birth satisfaction

Timepoint

12-24 h after childbirth

Method of measurement

Birth satisfaction scale-revised questionnaire

2**Description**

Pain intensity

Timepoint

About 30 min before the intervention and then 1, 2 and 3 h after starting the intervention

Method of measurement

Visual Analog Scale

3**Description**

Duration of the second stage of labor

Timepoint

After fetal expulsion

Method of measurement

Timer

4**Description**

Duration of the third stage of labor

Timepoint

After complete expulsion of the placenta

Method of measurement

Timer

5**Description**

Bishop Score

Timepoint

Just before the intervention and 2 h after starting the intervention

Method of measurement

Bishop Scoring Table

6**Description**

Hemoglobin and hematocrit

Timepoint

12- 24 h after delivery

Method of measurement

Laboratory test

7**Description**

Childbirth fear

Timepoint

At baseline and two h after starting intervention

Method of measurement

Delivery Fear Scale

8**Description**

Severity of postpartum fear

Timepoint

12-24 h and one Month after delivery

Method of measurement

Wijma Version B scale

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

Intervention group: Just after onset of active phase of labor (cervical dilation of 4-5 cm with regular uterine contractions), 10 mL of magnesium sulfate 50% made by Yara Teb Samen Pharmaceutical Company will be poured on the cervix through a 10 mL syringe during vaginal examination from the fingertips of the examiner, so that the whole cervix is impregnated with it. To better absorption of the drug, the women will be asked to sleep in bed for at least half an hour after pouring the drug. If the amniotic sac ruptures less than half an hour after pouring the drug, another 10 mL of magnesium sulfate will be poured on the cervix after discontinuing the discharge.

Category

Treatment - Drugs

2**Description**

Control group: Just after onset active phase of labor (cervical dilation of 4-5 cm with regular uterine contractions), 10 mL of placebo (distilled water) will be poured on the cervix through a 10 mL syringe during vaginal examination from the fingertips of the examiner, so that the whole cervix is impregnated with it. The women will be asked to sleep in bed for at least half an hour after pouring the placebo. If the amniotic sac ruptures less than half an hour after pouring the placebo, another 10 mL of distilled water will be poured on the cervix after discontinuing the discharge.

Category

Treatment - Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Taleghani Medical Research & Training Hospital

Full name of responsible person

Mansour Rezaei

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

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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

Tabriz 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

Tabriz University of Medical Sciences

Full name of responsible person

Sahar Ruhzende

Position

MSc student in midwifery

Latest degree

Bachelor

Other areas of specialty/work

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Person responsible for scientific inquiries

Contact

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Sakineh Mohammad-Alizadeh-charandabi

Position

Professor

Latest degree

Ph.D.

Other areas of specialty/work

Reproductive Health

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Person responsible for updating data

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Sharing plan

Deidentified Individual Participant Data Set (IPD)

Yes - There is a plan to make this available

Study Protocol

No - There is not a plan to make this available

Statistical Analysis Plan

No - There is not 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 deidentified IPD can be shared.

When the data will become available and for how long

Starting soon after publication of the study results.

To whom data/document is available

Data will be available for researchers working in academic institutions, as well as to chief editor and reviewers of the submitted manuscript.

Under which criteria data/document could be used

The data will be available to researchers upon request and submission of the proposal to perform meta-analysis using IPD. Also, in exceptional cases, data will be made available to chief-editor of the journals for checking.

From where data/document is obtainable

Refer to the email addresses (alizades@tbzmed.ac.ir).

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

The requests should be sent by email and data will be available within two week.

Comments