

A randomised trial to evaluate misoprostol for induction of labour following prelabour rupture of the amniotic membranes

Submission date
21/07/2005

Recruitment status
No longer recruiting

☐ Prospectively registered

☐ Protocol

Registration date
09/09/2005

Overall study status
Completed

☐ Statistical analysis plan

☒ Results

Last Edited
06/10/2009

Condition category
Pregnancy and Childbirth

☐ Individual participant data

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

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Contact details

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Additional identifiers

Protocol serial number

LWH315

Study information

Scientific Title

Acronym

The PROMMIS Trial - Prelabour Rupture Of the Membranes and MISoprostol

Study objectives

Vaginal misoprostol for cervical priming followed by titrated oral misoprostol is as safe and effective as dinoprostone and/or oxytocin for induction of labour in the presence of prelabour rupture of membranes (PROM)

Ethics approval required

Old ethics approval format

Ethics approval(s)

Not provided at time of registration

Primary study design

Interventional

Study design

Randomised active controlled parallel group trial

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Induction of labour

Interventions

Randomisation to dinoprostone or misoprostol

Intervention Type

Drug

Phase

Not Specified

Drug/device/biological/vaccine name(s)

Misoprostol, dinoprostone, oxytocin

Primary outcome(s)

1. Caesarean section (measure of safety)
2. Vaginal delivery within 24 hours of trial entry (measure of clinical effectiveness)

Key secondary outcome(s)

1. Labour
2. Delivery
3. Puerperium
4. Neonatal Morbidity
5. Satisfaction - Women's satisfaction, Caregiver's satisfaction
6. Womens' and midwives' views and experiences of participating in the PROMMIS Trial

Completion date

31/12/2005

Eligibility

Key inclusion criteria

1. Decision to induce labour in the presence of PROM
2. >34 weeks gestation
3. Singleton, live fetus
4. Normal admission cardiotocograph (CTG)

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

Female

Key exclusion criteria

1. Multiple pregnancy
2. Breech presentation
3. Previous Caesarean section

Date of first enrolment

01/02/2002

Date of final enrolment

31/12/2005

Locations

Countries of recruitment

United Kingdom

England

Study participating centre

Liverpool Women's Hospital NHS Foundation Trust

Liverpool

United Kingdom

L8 7SS

Sponsor information

Organisation

Liverpool Women's Hospital NHS Foundation Trust (UK)

ROR

<https://ror.org/04q5r0746>

Funder(s)

Funder type

Government

Funder Name

Liverpool Women's Hospital NHS Foundation Trust (UK)

Funder Name

NHS Support Funds

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/11/2008		Yes	No