# Misoprostol for treating postpartum haemorrhage: a randomised controlled trial

Submission date Recruitment status Prospectively registered 19/07/2004 No longer recruiting [ ] Protocol [ ] Statistical analysis plan Registration date Overall study status 20/07/2004 Completed [X] Results [ ] Individual participant data **Last Edited** Condition category 10/08/2007 Pregnancy and Childbirth

#### Plain English summary of protocol

Not provided at time of registration

#### Contact information

#### Type(s)

Scientific

#### Contact name

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#### Additional identifiers

Protocol serial number CRHS010215a

# Study information

Scientific Title

#### **Study objectives**

Not provided at time of registration

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Not provided at time of registration

#### Study design

Randomised controlled trial

#### Primary study design

Interventional

#### Study type(s)

Screening

#### Health condition(s) or problem(s) studied

Postpartum haemorrhage

#### **Interventions**

- 1. Intervention group: Misoprostol 200 µg tablets. Each woman receives 1 tablet orally, 2 tablets sublingually and 2 tablets rectally.
- 2. Control group: identical placebo tablets. Each woman receives 1 tablet orally, 2 tablets sublingually and 2 tablets rectally.

(All women receive routine accepted treatment for postpartum haemorrhage)

#### Intervention Type

Drug

#### Phase

**Not Specified** 

#### Drug/device/biological/vaccine name(s)

Misoprostol

#### Primary outcome(s)

Not provided at time of registration

#### Key secondary outcome(s))

Not provided at time of registration

#### Completion date

01/12/2003

## **Eligibility**

#### Key inclusion criteria

Women with postpartum haemorrhage defined as vaginal bleeding after childbirth considered to be excessive, and considered likely to be due to inadequate uterine contraction

#### Participant type(s)

#### **Patient**

#### Healthy volunteers allowed

No

#### Age group

Adult

#### Sex

**Female** 

#### Key exclusion criteria

Not provided at time of registration

#### Date of first enrolment

01/01/2002

#### Date of final enrolment

01/12/2003

### Locations

#### Countries of recruitment

South Africa

# Study participating centre P Bag X9047

East London South Africa 5201

# Sponsor information

#### Organisation

University of the Witwatersrand (South Africa)

#### **ROR**

https://ror.org/03rp50x72

# Funder(s)

#### Funder type

University/education

#### Funder Name

University of the Witwatersrand (South Africa) - No external funding; this trial was funded from the existing research unit budget, using full-time research staff.

# **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 | 06/08/2004   |            | Yes            | No              |