

What shall we do with the postdates woman who is not in labour?

Submission date	Recruitment status	<input type="checkbox"/> Prospectively registered
12/09/2003	No longer recruiting	<input type="checkbox"/> Protocol
Registration date	Overall study status	<input type="checkbox"/> Statistical analysis plan
12/09/2003	Completed	<input type="checkbox"/> Results
Last Edited	Condition category	<input type="checkbox"/> Individual participant data
02/06/2017	Pregnancy and Childbirth	<input type="checkbox"/> Record updated in last year

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

N0024112467

Study information

Scientific Title

What shall we do with the postdates woman who is not in labour?

Study objectives

Can we reduce the induction to delivery interval and Caesarean section rate in postdates primigravida women undergoing induction of labour with unfavourable cervix?

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)

Treatment

Health condition(s) or problem(s) studied

Pregnancy and Childbirth: Induction of labour

Interventions

1. Mifepristone
2. Standard care

Intervention Type

Drug

Phase

Not Applicable

Drug/device/biological/vaccine name(s)

Mifepristone

Primary outcome(s)

1. Length of time from induction start (first Prostin) to:
 - 1.1. Onset of labour
 - 1.2. Second stage
 - 1.3. Delivery
- 1.4. Timing and type of intervention (Caesarian Section, Forceps, Ventouse), if any
2. Antepartum haemorrhage (APH), postpartum haemorrhage (PPH) and other complications
3. Neonatal weight, cord pH and admission to Special Care Baby Unit (SCBU)

Key secondary outcome(s)

Not provided at time of registration

Completion date

31/12/2003

Eligibility

Key inclusion criteria

100 pregnant primiparous women who have not laboured at or beyond 41 weeks of gestation with cervical length >3 cm.

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

01/05/2002

Date of final enrolment

31/12/2003

Locations

Countries of recruitment

United Kingdom

England

Study participating centre

Fetal Medicine Unit

London

United Kingdom

E9 6SR

Sponsor information

Organisation

Department of Health (UK)

Funder(s)

Funder type

Government

Funder Name

Homerton University Hospital NHS Trust (UK)

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration