Impact of dalteparin in pregnancy

Submission date Prospectively registered Recruitment status 18/04/2008 No longer recruiting [] Protocol [] Statistical analysis plan Registration date Overall study status 29/04/2008 Completed [X] Results [] Individual participant data Last Edited Condition category 30/12/2020 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

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers

CER Sainte-Justine 1006

Study information

Scientific Title

Dalteparin in prevention of recurrence of severe obstetrical complications in women without thrombophilia

Study objectives

A prophylactic dose of dalteparin may prevent the recurrence of placental mediated complications in women without thrombophilia.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethics approval received from the Comité déthique de recherche (CER) du CHU Sainte-Justine on the 16th March 2000 (ref: 1006).

Study design

Randomised open trial

Primary study design

Interventional

Secondary study design

Randomised controlled trial

Study setting(s)

Hospital

Study type(s)

Treatment

Participant information sheet

Health condition(s) or problem(s) studied

Severe obstetrical complications

Interventions

Randomisation (1:1) to dalteparin (5000 IU daily) versus no treatment from randomisation until 36 - 37 weeks of pregnancy. Total duration of follow-up for all treatment arms will be from randomisation until departure from the hospital after delivery.

Intervention Type

Drug

Phase

Not Specified

Drug/device/biological/vaccine name(s)

Dalteparin

Primary outcome measure

Composite outcome of one of:

- 1. Severe preeclampsia
- 2. Severe IUGR
- 3. Major abruptio placentae

Primary outcomes will be measured at delivery.

Secondary outcome measures

- 1. Non-severe preeclampsia, measured at delivery
- 2. Non-severe IUGR, measured at delivery
- 3. Gestational age at delivery, measured at delivery
- 4. Pregnancy complications, measured at delivery
- 5. Hospitalisation length, measured at delivery and departure from the hospital after delivery
- 6. Admission to intensive care unit, measured at departure from the hospital after delivery

Overall study start date

16/03/2000

Completion date

20/06/2007

Eligibility

Key inclusion criteria

- 1. Normal thrombotic screen
- 2. Female patients aged more than 17 years
- 3. Pregnancy less than 17 weeks
- 4. Signature of the consent form
- 5. One of the following in previous pregnancy:
- 5.1. Severe preeclampsia with delivery less than 34 weeks
- 5.2. Severe intrauterine growth restriction (IUGR)
- 5.3. Abruptio placentae with delivery less than 34 weeks or foetal death
- 5.4. One idiopathic foetal death after 20 weeks
- 5.5. Two or more of idiopathic foetal death between 12 19 weeks

Participant type(s)

Patient

Age group

Adult

Sex

Female

Target number of participants

276

Total final enrolment

116

Key exclusion criteria

- 1. Multiple pregnancy
- 2. Alcohol or illicit drug use
- 3. Severe medical condition other than chronic hypertension
- 4. Foetal malformation or chromosomal anomaly
- 5. Uterine malformation or infection
- 6. Abdominal trauma
- 7. Any known cause of foetal death

Date of first enrolment

16/03/2000

Date of final enrolment

20/06/2007

Locations

Countries of recruitment

Canada

Study participating centre 3175 Côte sainte-Catherine

Montreal (QC) Canada H3T 1C5

Sponsor information

Organisation

Canadian Foundation for Women's Health (Canada)

Sponsor details

780 Echo Drive Ottawa (ON) Canada K1S 5R7 +1 800 561 2416 helpdesk@sogc.com

Sponsor type

Research organisation

Website

http://www.cfwh.org

ROR

https://ror.org/0278syk25

Funder(s)

Funder type

Research organisation

Funder Name

Canadian Foundation for Women's Health (Canada)

Funder Name

Pharmacia and Upjohn Inc. (USA)

Results and Publications

Publication and dissemination plan

Not provided at time of registration

Intention to publish date

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	01/01/2009	30/12/2020	Yes	No