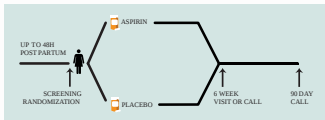


Pilot **PARTUM** Trial: Postpartum **A**spirin to **R**educe **T**hromboembolism **U**ndue **M**orbidity

Postpartum women at modest risk of VTE (deep vein thrombosis or pulmonary embolism) will be randomized to low-dose aspirin daily versus placebo daily for 6 weeks postpartum. Participation is simple:



Thank you for supporting the pilot PARTUM trial!

partumtrial.ca

OMNI Research Group:
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INCLUSION

ONE
of:

1. Mild inherited thrombophilia:
Heterozygous FVL or Heterozygous PGM or PC def or PS def
2. Bedrest (90%) for ≥ 7 days during pregnancy

OR

TWO
of:

3. Pre-pregnancy BMI ≥ 30 kg/m²
4. Smoking in past 12 months
5. Past superficial vein thrombosis
6. Pre-eclampsia
7. Stillbirth in current pregnancy (>20 weeks)
8. SGA infant <3 rd percentile
9. Unplanned cesarean delivery
10. Postpartum infection
11. Postpartum hemorrhage



Subjects are excluded if they are >48 hrs after delivery, received >2 LMWH doses or have had a prior VTE. See poster for full eligibility criteria.