

LOCAL OPERATING PROCEDURE

Approved by Quality & Patient Care Committee 21 June 2018

POSTPARTUM HAEMORRHAGE (PPH) – PREVENTION AND MANAGEMENT

This LOP is developed to guide clinical practice at the Royal Hospital for Women. Individual patient circumstances may mean that practice diverges from this LOP.

1. AIM

Early recognition and prompt appropriate intervention to minimise the impact of postpartum haemorrhage (PPH)

2. PATIENT

• A woman whose blood loss at or after childbirth is measured or estimated at ≥500mls, or who experiences hemodynamic compromise as a result of postpartum bleeding

3. STAFF

• Medical, nursing and midwifery staff

4. EQUIPMENT

- Two large bore intravenous (IV) cannulae (14–16 gauge)
- Blood tubes (pink, purple +/- blue topped)
- IV Starter Kit
- · Gloves
- Sphygmomanometer
- Personal protective equipment (PPE)
- Measuring equipment e.g. scales, jug, kidney dish
- Indwelling urinary catheter (IDC)
- PPH Box

5. CLINICAL PRACTICE

Prevention of PPH

- · Recommend active management of third stage of labour to each woman antenatally
- Consider additional prophylaxis for prevention of PPH for high risk woman (Appendix 1)

Treatment of PPH immediate management

- · Call for help
- · Activate Patient with Acute Condition for Escalation (PACE) call according to criteria
- Perform stepwise management of PPH as per flowchart (Appendix 2)
- · Identify underlying cause of PPH and check placenta and membranes are complete
- Replace volume by infusing warm crystalloid solution at least three times the measured volume of blood lost. Consult anaesthetic team if more than two litres crystalloid solution is required
- Consider treatment with uterotonic medications and/or intravenous (IV) tranexamic acid (Appendix 3)
- Keep the woman warm and administer high flow oxygen via facial mask
- Notify consultant obstetrician to attend if PPH > 1.5L and ongoing bleeding

Management of ongoing bleeding

- Escalate further as required e.g. PACE 2, code blue, consultant obstetrician attendance
- Transfer to theatre
- Utilise ROTEM to guide blood product replacement, led by anaesthetic team
- Activate Critical Bleeding Protocol (CBP) if criteria met, led by anaesthetic team. This can be used with or without ROTEM





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Postnatally

- Document estimated blood loss and treatments used for PPH
- Debrief woman and her family members/support people
- Debrief staff

6. DOCUMENTATION

- Medical Record
- Obstetric database
- PACE Notification
- IV Fluid Chart
- Fluid Balance Chart

7. EDUCATIONAL NOTES

- · Primary PPH is within 24 hours of birth
- · Secondary PPH is 24 hours to six weeks postpartum
- · Severe PPH is defined as blood loss of 1000 ml or more after childbirth
- Blood loss of <u>>2000ml</u> carries a significant risk for coagulopathy, and additional escalation is recommended when blood loss is more than this or if there is hemodynamic compromise
- Primary Prophylaxis/Active management of third stage. Routine prophylactic oxytocin administered after delivery of the anterior shoulder reduces the risk of PPH by more than 40% and is the most effective means of preventing PPH from uterine atony and is not associated with an increased risk of retained placenta. Active management of third stage involves:
 - o oxytocin
 - cord clamping and cutting
 - controlled cord traction (CCT)
- Aetiology:
 - TONE 70% of PPHs are caused by abnormalities of uterine contraction (atony)
 - TRAUMA 20% of PPHs are genital tract trauma
 - TISSUE 10% of PPHs are caused because placental or membrane tissue is retained
 - THROMBIN<1% of PPHs are caused by coagulation abnormalities. Abnormalities of coagulation may be present prior to or during pregnancy or may reflect the severity of blood loss during PPH
- When blood loss continues or woman is haemodynamically unstable, other less common causes need to be considered:
 - uterine inversion
 - uterine rupture
 - broad ligament haematoma
- PPH boxes are located in Delivery Suite, Birth Centre, Operating Theatre and both Postnatal Wards
- ROTEM is a point of care whole blood haemostasis testing method
- CBP replaced Massive Transfusion Protocol (MTP) in April 2018
- Uterine/vaginal tamponade may be undertaken by the use of rolled gauze or intrauterine cavity balloon
- Misoprostol, a prostaglandin E1 analogue, is not currently recommended for routine prevention and control of PPH. Its use is unlicensed, however, it may be used as an adjunct to other medications in cases of severe PPH.
- Tranexamic acid has been used to treat PPH. In a meta-analysis (two trials (20,412 women)) it was found that IV tranexamic acid reduces the risk of maternal death due to bleeding (risk ratio (RR) 0.81, 95% confidence interval (CI) 0.65 to 1.00; two trials, 20,172 women; quality of evidence: moderate). The effect was more evident in women given treatment between one and three hours after giving birth with no apparent reduction when given after three hours.





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8. RELATED POLICIES / PROCEDURES / CLINICAL PRACTICE LOP

- Critical Bleeding Protocol (CBP) Business Rule. POWH CLIN072
- Third Stage Management Following Vaginal Birth
- Blood Products Management of Pregnant Woman Unable to Use Blood Products
- Patient with Acute Condition for Escalation (PACE): Management of the Deteriorating ADULT and MATERNITY Inpatient. SESLHDPR/283
- NSW Health Policy Directive. Maternity Prevention, Early Recognition & Management of Postpartum Haemorrhage (PPH) 2017. GL 2017_018
- NSW Health Policy Directive PD2007_040 Open Disclosure
- NSW Health Policy Directive PD2007_061 Incident Management
- Balloon Placement for Uterine Tamponade
- · Perineal/Genital Tract Repair
- · Labelling of Injectable Medicines, Fluids, and Lines
- Maternal Collapse
- Escalation for Birthing Services
- Maternity Prevention, Detection, Escalation and Management of Postpartum Haemorrhage (PPH) GL2017_018

9. RISK RATING

High

10. NATIONAL STANDARD

• CC – Comprehensive Care

11. REFERENCES

- 1. RCOG 2016. Postpartum Haemorrhage Prevention and Management. Green-Top Guideline No. 52
- 2. Queensland Maternity and Neonatal Clinical Guidelines Program. 2018 Primary postpartum haemorrhage MN18.1-V7-R23
- 3. Pairman S, Tracy S, Thorgood C and Pincombe V. Midwifery Preparation for practice 2010
- 4. Mousa HA, Blum J, Abou El Senoun G, Shakur H, Alfirevic Z. Treatment for primary postpartum haemorrhage. Cochrane Database Systemic Reviews. 2014 Issue 2 Feb 13;(2):CD003249.
- 5. Shakur H, Beaumont D, Pavord S, Gayet-Ageron A, Ker K, Mousa HA. Antifibrinolytic drugs for treating primary postpartum haemorrhage. Cochrane Database Systemic Reviews 2018 Feb 20;2:CD012964.
- 6. Effect of early tranexamic acid administration on mortality, hysterectomy, and other morbidities in women with post-partum haemorrhage (WOMAN): an international, randomised, double-blind, placebocontrolled trial. Shakur, Haleema et al. The Lancet, Volume 389, Issue 10084, 2105 - 2116

REVISION & APPROVAL HISTORY

Reviewed and endorsed Maternity Services LOPs 19/6/18

Approved Quality & Patient Care Committee 4/2/16

Reviewed and endorsed Maternity Services LOPs group December 2015

Approved Quality & Patient Safety Committee December 2012

Amendment to dosages in appendix May 2014

Reviewed and endorsed Maternity Services LOPs group December 2012

Reviewed Obstetric Clinical Guidelines Group Sept 2010 – Approved Quality & Patient Safety Committee 21/10/10

Reviewed July 2007 – Approved Clinical Performance & Quality Committee August 2007 Endorsed Maternity Services Clinical Committee 10/12/02 – Approved Quality Council 16/12/02

FOR REVIEW: JANUARY 2019

3.

APPENDIX1

RISK FACTORS FOR PPH REQUIRING ADDITIONAL PROPHYLAXIS:

- EITHER ERGOMETRINE (IF NO CONTRAINDICATIONS) 250mcg IM/IV
- <u>AND/OR OXYTOCIN INFUSION (40 UNITS OXYTOCIN (SYNTOCINON) IN 1000MLS SODIUM</u> <u>CHLORIDE 0.9% @ 250mLs/hr)</u>

SUSPECTED OR PROVEN PLACENTAL ABRUPTION	
MULTIPLE PREGNANCY	
RETAINED PLACENTA >30 MINUTES	
PRE ECLAMPSIA/GESTATIONAL HYPERTENSION	
BIRTH BY EMERGENCY CAESAREAN SECTION	
PREVIOUS PPH	
OPERATIVE VAGINAL BIRTH/SHOULDER DYSTOCIA	
PROLONGED LABOUR>12 HOURS	
SECOND STAGE OF LABOUR>2 HOURS	
VON WILLEBRAND'S DISEASE	
ANAEMIA (<9 g/L)	
GRAND MULTIPARITY	

<u>OTHER RISK FACTORS (CONSIDER ADDITIONAL PROPHYLAXIS, PARTICULARLY IN</u> <u>THE CASE OF MULTIPLE RISK FACTORS)</u>

ASIAN ETHNICITY	
OBESITY (BMI>30)	
INDUCTION/AUGMENTATION OF LABOUR	
BABY WEIGHT>4 KG	
PYREXIA IN LABOUR	
AGE >40 YEARS	
PRECIPITATE LABOUR	
MULTIPLE OR LARGE FIBROIDS	
POLYHYDRAMNIOS	

A P P E N D I X 2 FLOWCHART – PRIMARY POSTPARTUM HAEMORRHAGE ≥ 500ML

IMMEDIATE MANAGEMENT – STEPS MAY OCCUR CONCURRENTLY

- Call for help, initiate PACE according to criteria, ensure neonatal safety
- Lie woman flat, massage fundus (expel clots if indicated) and provide reassurance
- Commence oxygen by facial mask
- Ensure 10 units of oxytocin intramuscularly (IM) has been given
- Insert two large bore cannulae (14g or 16g), send blood for FBC, Group and hold +/- cross-match, coags, biochemistry
- Commence volume replacement, ideally with warm crystalloid
- Monitor blood pressure, pulse, respiration, SpO² every 5 minutes and temperature every 15 minutes
- Keep woman warm
- Insert IDC



APPENDIX 3

MEDICATIONS T	O USE WITH PPH	CONTRAINDICATIONS/CAUTIONS
1 st Line Treatment	 Ergometrine Give 250 microgram either IM or slow IV infusion with antiemetic. This can be repeated if required. Onset of action: IM is 5-7 minutes, lasts 3 hours IV has rapid onset within 1 minute and lasts 45 minutes 	Contraindications: • ergot alkaloid hypersensitivity • retained placenta • pre-eclampsia/eclampsia • sepsis • peripheral vascular disease • heart disease • current or past history of hypertension • impaired hepatic/renal function
2 ND LINE TREATMENT	 Oxytocin Infusion Add 40 units to 1 litre of Normal Saline (sodium chloride 0.9%) and run at 250 mLs/hour via infusion pump Onset of action: IV < 1 minute, lasts <30 minutes. IM 2-4 minutes, lasts 30-60 minutes 	Contraindication – known hypersensitivity
3 RD Line Treatment	 Misoprostol Give 800 micrograms rectally Onset of action per rectum has slow uptake (100 minutes) but prolonged duration (4 hours). Off label use 	 Contraindication – known hypersensitivity Caution - asthma. Side effects: diarrhoea abdominal pain shivering/fever
4 th Line Treatment	 Tranexamic acid Give as a slow IV push 1gm/10mLs over 10 minutes (1mL per minute) If required, follow 30 minutes later with infusion of 1g diluted in sodium chloride or glucose solutions 500mls, at 250mLs/hour via infusion pump 	 Contraindications: Active thromboembolism including deep vein thromboses, pulmonary embolus, cerebral thrombosis thrombosis risk, including family history (unless anticoagulated) acquired colour vision disturbance subarachnoid haemorrhage Caution in renal impairment Side effects - dizziness and hypotension
5 TH Line Treatment	 Prostaglandin F2 Alpha Ensure an IV line, cardiac monitoring and O² therapy are in place before administration An anaesthetist should be in attendance Dilute 5mg (1mL) of Prostaglandin F2 Alpha with 9 ml of Normal Saline to equal 10mLs volume Discard 4mL to leave 6mL = 3mg or 500 microgram/mL Give 2 mL (or maximum 1 mg at a time) by a medical officer injecting into the uterine myometrium with the 22G Spinal Needle (BD®) 	 Caution: asthma hypertension active cardiac, renal or hepatic disease known hypersensitivity Side effects: nausea bronchospasm vomiting/diarrhoea headache flushing/pyrexia uterine rupture cardiac arrest
OR 5 TH LINE TREATMENT	 Carboprost[®] Ensure an IV line, cardiac monitoring and oxygen therapy are in place before administration An anaesthetist should be in attendance Give 250 microgram (1mL) via IM or intramyometrial injection. Intramyometrial injection is an 'off label' route of administration and therefore must be administered by a medical officer Once in OT, the dose can be repeated as required every 15-90 minutes to a maximum of 2mg (8 doses). This medication is imported from overseas via the Special Access Scheme (SAS). Please complete an SAS form and return to Pharmacy. Where possible, obtain consent from patient and document in clinical notes. 	 Contraindications: acute pelvic inflammatory disease cardiac/pulmonary/renal/hepatic disease known hypersensitivity to prostaglandin Cautions: asthma anaemia diabetes epilepsy hyper/hypotension jaundice uterine surgery Side effects: hypertensive crisis fever with rigors headache paraesthesia diarrhoea, nausea and vomiting breast tenderness dystonia pulmonary oedema