Management of Postpartum Haemorrhage (PPH)

Objectives: To provide advice on the management of postpartum haemorrhage.

Outcomes: Minimising risks for the patient associated with Postpartum Haemorrhage.

Target audience: All health practitioners providing maternity care and patients.

Values: The evidence was reviewed by the Women’s Health Committee (RANZCOG), and applied to local factors relating to Australia and New Zealand.

Background: This statement was first developed by Women’s Health Committee in March 2011 and the last full review was conducted in March 2014, July 2017. Minor amendments were made in May 2015 and February 2016.

Funding: The development and review of this statement was funded by RANZCOG.

This statement has been developed and reviewed by the Women’s Health Committee and approved by the RANZCOG Board and Council.

A list of Women’s Health Committee Members can be found in Appendix B.

Disclosure statements have been received from all members of this committee.

Disclaimer This information is intended to provide general advice to practitioners. This information should not be relied on as a substitute for proper assessment with respect to the particular circumstances of each case and the needs of any patient. This document reflects emerging clinical and scientific advances as of the date issued and is subject to change. The document has been prepared having regard to general circumstances.

First endorsed by RANZCOG: March 2011
Current: July 2017
Review due: July 2020
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1. **Patient Summary**

Heavy bleeding after a baby is born (postpartum haemorrhage) is a complication of pregnancy that has the potential to be very serious, even resulting in death in rare cases. Some women will have risk factors for heavy bleeding, but most will not. This means that it is very difficult to predict when heavy bleeding might occur and essentially all women giving birth are at risk of having a severe postpartum haemorrhage.

The use of some simple measures — receiving a small dose of medication to help the uterus contract after birth and assisting with delivering the placenta — all reduce the risk of heavy bleeding.

When heavy bleeding occurs, the birth attendants must act quickly and use proven techniques to stop the bleeding and help the woman recover. Because postpartum haemorrhage occurs so unpredictably, all women giving birth should have timely access to skilled care and all the resources needed for managing heavy bleeding. This is one of many reasons why the College recommends that all women birth in (or adjacent to) an appropriately equipped hospital.

2. **Summary of recommendations**

<table>
<thead>
<tr>
<th>Recommendation 1</th>
<th>Grade and reference</th>
</tr>
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<tbody>
<tr>
<td>Active management of the third stage of labour (administration of prophylactic oxytocics and assisting delivery of the placenta) should be recommended to all pregnant women as this reduces the risk of PPH and the need for blood transfusion. Prophylactic oxytocics should be recommended for the management of the third stage of labour, whether following vaginal or caesarean birth, as they reduce the risk of PPH by at approximately 50%.</td>
<td>Evidence-based recommendation</td>
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<thead>
<tr>
<th>Recommendation 2</th>
<th>Grade and reference</th>
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<tr>
<td>All those managing births regardless of location or complexity, must do so with a clear guideline for the management of a Postpartum Haemorrhage should it occur. The guideline should include acute management at the place of birth and the procedure for timely escalation should the haemorrhage not respond rapidly to initial measures.</td>
<td>Consensus-based recommendation</td>
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3. **Introduction**

Postpartum haemorrhage (PPH) remains a major cause of both maternal mortality and morbidity within Australia and New Zealand. PPH is common, with an incidence in Australia of between five and fifteen percent. While the majority of these cases are minor, requiring little active management and causing minimal morbidity, it must be remembered that PPH remains a leading cause of maternal death both globally and in Australia and New Zealand.

**Definitions**

Traditionally PPH has been defined as a blood loss of 500ml or more during puerperium and severe PPH as a blood loss of 1000ml or more. Further classification of PPH into primary (within 24 hours of delivery) and secondary (between 24 hours and six weeks postpartum) is also well established.

**Background Physiology**

In order to prevent and treat PPH, the clinician requires an understanding of the physiological mechanisms that arrest the blood flow to the placental bed following placental separation. The blood flow to the placental bed varies with gestation but is approximately 750 ml/min at term. Given that maternal blood volume may only be approximately 7 litres and blood loss may be life-threatening with unreplace...
loss of as little as 30%, it can be seen that a mother can become critically unwell in just a few minutes if blood flow to the placental bed is not rapidly arrested with placental separation.

Aetiology

The principle physiological mechanism for avoiding excessive blood loss is constriction of the blood vessels supplying the placenta bed by uterine contraction. This is supplemented by common haemostasis: vasoconstriction, platelet aggregation and clot formation. A classification of the causes of PPH are listed in table 1.

Table 1. Aetiology of PPH

Traumatic Site ("Trauma")
- Uterine rupture, Cervical Tear, Vaginal Tear, Perineal Tear

Placental Site
- Uterine Atony ("Tone")
  - Prolonged labour and particularly prolonged 2nd stage of labour
  - Increasing Parity
  - Oxytocin withdrawal
  - Uterine overdistension
    - Multiple pregnancy, Polyhydramnios, Macrosomia
  - Instrumental Birth
- Retained Products of Conception ("Tissue")
- Placenta Praevia
- Placenta Accreta
- Uterine Inversion
- Bleeding Disorder
  - Thrombocytopenia
  - Disseminated Intravascular Coagulation ("Thrombin")
    - Severe Preeclampsia, Placental Abruption, Sepsis, FDIU, Amniotic Fluid Embolism
  - Hereditary Bleeding Disorders e.g. von Willebrand’s
4. Discussion and recommendations

4.1 How can postpartum haemorrhage be prevented?

4.1.1 What are the risk factors?
A large number of risk factors for PPH have been identified but most cases of PPH have no identifiable risk factor. For those women known to have risk factors for PPH appropriate management should be instigated in both the antenatal and intrapartum periods to mitigate this risk. All women should birth in a unit with rapid access to blood and blood products and have antenatal correction of anaemia. Where a woman is likely to decline the administration of blood products, risk of death from postpartum haemorrhage becomes considerably greater and prophylaxis even more important than otherwise.

4.1.2 What are important considerations in the event of abnormal placentation?
All women must have placental location determined by antenatal ultrasound. Appropriate recognition, preparation and management of women with placenta praevia or suspected morbidly adherent placentation is crucial as these conditions are associated with increased risk of catastrophic haemorrhage and maternal mortality.

4.1.3 How should the third stage of labour be managed?
Active management of the third stage of labour (use of prophylactic oxytocics and assisting delivery of the placenta) should be practised as this reduces the risk of PPH and the need for blood transfusion. Prophylactic oxytocics should be used for the management of the third stage of labour, whether following vaginal or caesarean birth, as they reduce the risk of PPH by at least 50%. While misoprostol has been used in routine management of the third stage of labour, quality trials in the hospital setting have reported that it is less effective than oxytocin and is associated with a greater incidence of side effects. Oxytocin should remain the drug of choice for this indication and misoprostol can be an alternative when the use of oxytocin is not possible. Women may occasionally request physiological management of the third stage without the use of an oxytocic. It is important that these women are adequately informed of the increased risks of bleeding associated with this practice.

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<td>Evidence-based recommendation 1</td>
</tr>
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4.2 What are the management considerations for Postpartum Haemorrhage?
Effective team management of PPH involves recognition, communication, resuscitation, monitoring and investigation and directed treatment. Once a PPH has been recognised these components should be conducted simultaneously for optimal patient care. Some guidelines invoke basic measures for estimated blood loss (EBL) of 500ml-1000ml with no clinical shock and a full protocol of measures for EBL greater than 1000ml and continuing bleeding, or where there is evidence of clinical shock. It is important to consider both the patient’s prior haemoglobin and her total blood volume when assessing the severity of PPH. Total blood volume at term is approximately 100ml/kg (i.e. 7000ml for a 70 kg woman, but only 5000ml for a 50kg woman).
1. Recognition

Assessment of ongoing blood loss is an essential aspect of post-partum care. Visual estimation of blood loss is notoriously unreliable and often underestimates true blood loss. Weighing drapes, pads and swabs will be more accurate but it should be remembered that some amniotic fluid will be included in the estimation. Clinical signs of shock or tachycardia should prompt a thorough assessment of the mother including an accurate appraisal of blood loss, both concealed and revealed.

2. Communication

The successful management of PPH requires a multidisciplinary team approach. The clinical team involved, their response to PPH, and ability to escalate this response in the face of severe haemorrhage will vary according to the setting and circumstance of delivery. All centres providing obstetric care should implement and regularly review a clear plan of communication, resuscitation and directed treatment to respond to PPH. Senior obstetric and midwifery staff will be needed in the first instance, but other clinicians (such as anaesthetists, haematologists or transfusion specialists, intensivists and sub-specialty surgeons) may be called upon in the setting of more serious haemorrhage. In all cases of PPH, a second clinician (usually the anaesthetist) is vital in ensuring adequate resuscitation, especially while the obstetrician is busy instituting operative management. Often haematologists are required to co-ordinate the transfusion of blood products. Communication with the patient and their support person is important as this can be a frightening event for all involved.

3. Resuscitation

The cornerstone of resuscitation is restoration of blood volume and oxygen-carrying capacity. A simple ‘ABC’ approach is often used initially but clinical judgement should be used to guide resuscitation in each situation.  

- Immediately call for help  
- Rapidly assess for danger to self and others  
- Assessment of airway and breathing initially with administration of high flow oxygen is recommended.  
- Wide-bore intravenous access should be established with blood sent for full blood count, coagulation profile and cross-match.  
- Rapid infusion with fluids, ideally warmed, should be begun once intravenous access is achieved.  
- The use of group specific or group O Rh(D)-negative blood should be considered early to restore oxygen carrying capacity.  
- It is critical that facilities providing obstetric care have, and adhere to, a massive transfusion protocol with which all staff are familiar. Many larger hospitals will already have such a protocol in place, but a template can be found in Appendix A.  
- Additional measures such as keeping the woman warm and positioned flat are also important.

4. Monitoring and investigation

Appropriate monitoring and investigation should be guided by clinical judgement, but in all cases of PPH, should, at a minimum, include the recording of observations at regular intervals, (not monitoring and already done by now) and repeating, as indicated, in an appropriate time frame the haematological investigations. Pulse rate, blood pressure, oxygen saturation and urinary output measurement should be instigated for any significant (>1000ml) or ongoing PPH, and invasive monitoring of arterial blood pressure or central venous pressure may be necessary depending upon the clinical situation. Consideration must be given to the most appropriate place of care in women with severe PPH; this may be a high dependency care or intensive care unit in some instances. Where patients need to be transferred to a more highly equipped facility for management of PPH, the need for transfer should be anticipated and initiated early. In the
meantime, aggressive resuscitation should continue and regular communication with the receiving unit is essential.

Appropriate prophylaxis for venous thromboembolism should also be instituted once the acute situation has been controlled.

5. Management of PPH

Management of PPH invariably involves addressing the causes of bleeding, commonly known as ‘the four Ts’. A fifth ‘T’ has been added to emphasise the important role of theatre and surgery in managing all causes of PPH. There is little randomised control trial (RCT) evidence to guide the management of PPH. The only Cochrane review on primary PPH identified only three suitable trials; all concerned the use of misoprostol. There was inadequate evidence to comment on the utility of surgical, radiological, haemostatic or other pharmacological interventions. Hence most interventions are guided by best practice rather than high quality evidence.

a. Tone

Uterine atony is the most common cause of primary PPH but clinical assessment should be used to exclude other causes. The following interventions have all been used to stop the bleeding, generally in the stepwise progression as presented here.

i. Mechanical:
   • Uterine massage or bimanual uterine compression (emptying the bladder may assist in this process).

ii. Pharmacological:
   The following agents are useful in the management of PPH. They are commonly given in combination and, in the absence of individual contraindications, a patient may be given all four in the event of severe ongoing atonic bleeding. Because of the difficulties in undertaking clinical trials in the circumstances of unexpected PPH, the outcomes of the uterotonic in varying combinations to manage PPH have not been assessed by sufficiently-powered randomised controlled trials. However, their use is strongly recommended if the atonic haemorrhage is continuing.

   • Oxytocin (Syntocinon)
     o 5 units by slow intravenous injection (if already administered for 3rd stage management, a repeat dose may be given).
     o 40 units in an intravenous infusion over 4 hours
   • Ergometrine 0.25 mg by slow intravenous or intramuscular injection, repeated if necessary 5 minute up to a maximum of 1.0 mg; in the absence of contraindications.
   • Misoprostol (up to 1000mcg) rectally. Whilst many studies have studied the use of misoprostol to manage the third stage of labour, fewer have dealt with using misoprostol to treat PPH. In the hospital setting, there is evidence to suggest that misoprostol is clinically equivalent to further oxytocin in women who have already received prophylactic oxytocin when used for excessive post-partum bleeding due to suspected uterine atony. Prostaglandin and its analogues are the most potent of the uterotonic but also have the most serious adverse effect profile which includes severe hypertension and bronchospasm (therefore contraindicated where there is a significant history of asthma). In Australia, following the discontinuation of dinoprost trometamol, alternatives may be accessed through the TGA Special Access Scheme (SAS).
     o 15-methyl-PGF2α (carboprost; Prostinfenem) which may be administered in one of two ways:
       ▪ Intra-muscular injection of 0.25mg, in repeated doses as required at intervals of not less than 15 minutes to a maximum total cumulative dose of 2.0mg (ie up to 8 doses)
Management of Postpartum Haemorrhage

- Intramyometrial injection of 0.5mg, under the responsibility of the administering clinician as it is not recommended by the manufacturer for intramyometrial use.

The off-label use of these medications are considered routine for this indication with high quality evidence to support its use.

b. Trauma of the genital tract.

Thorough assessment of the entire genital tract is essential. The perineum, vagina and cervix should all be visually inspected for bleeding sources. Pressure should be applied to bleeding areas and repair attempted, either in the labour ward or the operating theatre if required.

If the patient is shocked and the amount of vaginal bleeding is normal, consider intra-abdominal sources such as ruptured uterus, broad ligament haematoma, subcapsular liver rupture, ruptured spleen, and ruptured splenic artery, hepatic artery or pancreatic artery aneurysm.

c. Tissue (retained products of conception)

This is usually due to retained placenta, cotyledon or membranes. If the placenta has been delivered assess for obvious missing tissue. If the placenta has not been delivered and cannot be delivered easily by controlled cord traction, empty the bladder and transfer the patient to theatre for manual removal of placenta.

Even when the placenta appears complete, there is still the possibility of unrecognised retention of a cotyledon or segment of a bipartite placenta. Hence, uterine exploration is indicated for all cases of persistent postpartum haemorrhage.

d. Thrombin (abnormalities of coagulation)

i) Tranexamic Acid
   The “WOMAN” Trial demonstrated efficacy of the administration of 1g of tranexamic acid intravenously with a clinical diagnosis of PPH. The dose was repeated after 30 minutes if bleeding was persistent. The authors concluded that: “Tranexamic acid reduces death due to bleeding in women with post-partum haemorrhage with no adverse effects. When used as a treatment for postpartum haemorrhage, tranexamic acid should be given as soon as possible after bleeding onset.”
   Tranexamic acid should only be administered in the context of overall patient management; the protocol should include strict attention to the control of bleeding, physiological and metabolic parameters, coagulation status and temperature maintenance.

ii) Replacement of Platelets and Clotting Factors
   - Early replacement of clotting factors and platelets is essential in the management of severe PPH.
   - “Point of Care” testing has an increasing role in the assessment of PPH (e.g. ROTEM).
   - Particular attention should be made to the fibrinogen level with replacement (cryoprecipitate or fibrinogen concentrate if available) where the fibrinogen is < 1.5-2g/L.

iii) Management of specific abnormalities such as Von Willebrand’s disease or severe thrombocytopenia is beyond the scope of this guideline.

e. Theatre

Surgical interventions should be initiated sooner rather than later, especially hysterectomy in cases of uterine rupture, placenta accreta or uncontrolled massive haemorrhage. The following is a list of some
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available procedures. This should not necessarily be a step-wise progression and both order and utilisation will depend on the services/clinical experience available and the individual clinical circumstances.

i. Balloon tamponade. Several case series have been published reporting the results of using a Foley catheter, Bakri balloon, Rusch balloon or Sengstaken-Blackmore oesophageal catheter with good results where the uterus is empty and contracting.3

ii. Haemostatic brace suturing (such as the B-Lynch suture).

iii. Bilateral ligation of uterine arteries.

iv. Bilateral ligation of internal iliac arteries by an experienced operator.

v. Selective arterial embolisation. This intervention can only be achieved in institutions with timely access to both radiological expertise and equipment. It is important to note that time delays in accessing embolisation can occur and should not preclude alternate surgical treatment.

vi. Hysterectomy.

4.3 How should secondary PPH be managed?

Secondary PPH is usually associated with endometritis (with or without retained products of conception). Conventional treatment usually includes antibiotic therapy and, uterotonics in some cases. In situations of excessive or continued bleeding surgical intervention, particularly the evacuation of retained products, should be considered, irrespective of ultrasound findings.1
5. References


6. Other suggested reading


New Zealand Ministry of Health National Consensus Guidelines on the Treatment of Postpartum Haemorrhage


7. **Links to other College statements**

Evidence-based Medicine, Obstetrics and Gynaecology (C-Gen 15)

8. **Patient information**

A range of RANZCOG Patient Information Pamphlets can be ordered via:
## Appendices

### Appendix A – Patient Blood Management Guidelines: Module 1, Massive transfusion protocol template

**Massive transfusion protocol (MTP) template**

The information below, developed by consensus, broadly covers areas that should be included in a local MTP. This template can be used to develop or MTP to meet the needs of the local institution’s patient population and resources.

1. **Senior clinician determines that patient meets criteria for MTP activation**
2. **Baseline:**
   - Full blood count, coagulation screen (PT, INR, APTT, fibrinogen), biochemistry, arterial blood gases
3. **Notify transfusion laboratory (insert contact no.) to:**
   - ‘Activate MTP’
4. **Laboratory staff**
   - Notify haematologist/transfusion specialist
   - Prepare and issue blood components as requested
   - Anticipate repeat testing and blood component requirements
   - Minimise test turnaround times
   - Consider staff resources
5. **Haematologist/transfusion specialist**
   - Liaise regularly with laboratory and clinical team
   - Assist in interpretation of results, and advise on blood component support

**Senior clinician**

- **Request:**
  - 4 units RBC
  - 2 units FFP
  - Consider:
    - 1 acute therapeutic dose platelets
    - tranexamic acid in trauma patients
    - Include:
      - cryoprecipitate if fibrinogen < 1 g/L
      - Or locally agreed configuration

**AIM FOR:**

- Temperature > 35°C
- pH > 7.2
- Base excess < −6
- Lactate < 4 mmol/L
- Ca²⁺ > 1.1 mmol/L
- Platelets > 50 × 10⁹/L
- PT/APTT < 1.5 × normal
- INR ≤ 1.5
- Fibrinogen > 1.0 g/L

**OPTIMISE:**

- Oxygenation
- Cardiac output
- Tissue perfusion
- Metabolic state

**MONITOR**

(every 30-60 mins):

- Full blood count
- Coagulation screen
- Ionised calcium
- Arterial blood gases

**Bleeding controlled?**

- **YES**
- **NO**

**Notify transfusion laboratory to:**

- ‘Cease MTP’
Suggested criteria for activation of MTP

- Actual or anticipated 4 units RBC in < 4 hrs, + haemodynamically unstable, +/- anticipated ongoing bleeding
- Severe thoracic, abdominal, pelvic or multiple long bone trauma
- Major obstetric, gastrointestinal or surgical bleeding

Initial management of bleeding
- Identify cause
- Initial measures:
  - compression
  - tourniquet
  - packing
- Surgical assessment:
  - early surgery or angiography to stop bleeding

Specific surgical considerations
- If significant physiological derangement, consider damage control surgery or angiography

Cell salvage
- Consider use of cell salvage where appropriate

Resuscitation
- Avoid hypothermia, institute active warming
- Avoid excessive crystalloid
- Tolerate permissive hypotension (BP 80–100 mmHg systolic) until active bleeding controlled
- Do not use haemoglobin alone as a transfusion trigger

Special clinical situations
- Warfarin:
  - add vitamin K, prothrombinex/FFP
- Obstetric haemorrhage:
  - early DIC often present, consider cryoprecipitate
- Head injury:
  - aim for platelet count > 100 x 10^9/L
  - permissive hypotension contraindicated

Dosage
- Platelet count < 50 x 10^9/L
- INR > 1.5
- Fibrinogen < 1.0 g/L
- Tranexamic acid

Loading dose: 1 g over 10 min, best duration 1 g over 6 h

Considerations for use of rFVIIa

- The routine use of rFVIIa in trauma patients is not recommended due to its lack of effect on mortality (Grade B) and variable effect on morbidity (Grade C). Institutions may choose to develop a process for the use of rFVIIa where there is:
  - uncontrolled haemorrhage in salvageable patient, and
  - failed surgical or radiological measures to control bleeding, and
  - adequate blood component replacement, and
  - pH > 7.2, temperature > 34°C.

Discuss dose with haematologist/transfusion specialist

*RFVIIa is not licensed for use in this situation; all use must be part of practice review.

<table>
<thead>
<tr>
<th>ADP</th>
<th>arteriole dilation</th>
<th>FFP</th>
<th>fresh frozen plasma</th>
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<tbody>
<tr>
<td>INR</td>
<td>international normalized ratio</td>
<td>BP</td>
<td>blood pressure</td>
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<tr>
<td>DIC</td>
<td>disseminated intravascular coagulation</td>
<td>PT</td>
<td>prothrombin time</td>
</tr>
<tr>
<td>RBC</td>
<td>red blood cell</td>
<td>rFVIIa</td>
<td>activated recombinant factor VII</td>
</tr>
<tr>
<td>APTT</td>
<td>activated partial thromboplastin time</td>
<td>MTP</td>
<td>massive transfusion protocol</td>
</tr>
<tr>
<td>FBC</td>
<td>full blood count</td>
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Appendix B Women’s Health Committee Membership

<table>
<thead>
<tr>
<th>Name</th>
<th>Position on Committee</th>
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<tbody>
<tr>
<td>Professor Yee Leung</td>
<td>Chair</td>
</tr>
<tr>
<td>Dr Joseph Sgroi</td>
<td>Deputy Chair, Gynaecology</td>
</tr>
<tr>
<td>Associate Professor Janet Vaughan</td>
<td>Deputy Chair, Obstetrics</td>
</tr>
<tr>
<td>Associate Professor Ian Pettigrew</td>
<td>EAC Representative</td>
</tr>
<tr>
<td>Dr Tal Jacobson</td>
<td>Member</td>
</tr>
<tr>
<td>Dr Ian Page</td>
<td>Member</td>
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<tr>
<td>Dr John Regan</td>
<td>Member</td>
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<tr>
<td>Dr Craig Skidmore</td>
<td>Member</td>
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<tr>
<td>Dr Lisa Hui</td>
<td>Member</td>
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<tr>
<td>Dr Bernadette White</td>
<td>Member</td>
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<tr>
<td>Dr Scott White</td>
<td>Member</td>
</tr>
<tr>
<td>Associate Professor Kirsten Black</td>
<td>Member</td>
</tr>
<tr>
<td>Dr Greg Fox</td>
<td>College Medical Officer</td>
</tr>
<tr>
<td>Dr Marilyn Clarke</td>
<td>Chair of the ATSI WHC</td>
</tr>
<tr>
<td>Dr Martin Byrne</td>
<td>GPOAC Representative</td>
</tr>
<tr>
<td>Ms Catherine Whitby</td>
<td>Community Representative</td>
</tr>
<tr>
<td>Ms Sherryn Elworthy</td>
<td>Midwifery Representative</td>
</tr>
<tr>
<td>Dr Amelia Ryan</td>
<td>Trainee Representative</td>
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</tbody>
</table>

Appendix C Overview of the development and review process for this statement

i. Steps in developing and updating this statement

This statement was originally developed in March 2011 and was most recently reviewed in March 2017. The Women’s Health Committee carried out the following steps in reviewing this statement:

- Declarations of interest were sought from all members prior to reviewing this statement.
- Structured clinical questions were developed and agreed upon.
- An updated literature search to answer the clinical questions was undertaken.
- At the March 2014 face-to-face committee meeting, the existing consensus-based recommendations were reviewed and updated (where appropriate) based on the available body of evidence and clinical expertise. Recommendations were graded as set out below in Appendix C part iii)

ii. Declaration of interest process and management

Declaring interests is essential in order to prevent any potential conflict between the private interests of members, and their duties as part of the Women’s Health Committee.

A declaration of interest form specific to guidelines and statements was developed by RANZCOG and approved by the RANZCOG Board in September 2012. The Women’s Health Committee members were required to declare their relevant interests in writing on this form prior to participating in the review of this statement.
Members were required to update their information as soon as they become aware of any changes to their interests and there was also a standing agenda item at each meeting where declarations of interest were called for and recorded as part of the meeting minutes.

There were no significant real or perceived conflicts of interest that required management during the process of updating this statement.

iii. Grading of recommendations

Each recommendation in this College statement is given an overall grade as per the table below, based on the National Health and Medical Research Council (NHMRC) Levels of Evidence and Grades of Recommendations for Developers of Guidelines. Where no robust evidence was available but there was sufficient consensus within the Women’s Health Committee, consensus-based recommendations were developed or existing ones updated and are identifiable as such. Consensus-based recommendations were agreed to by the entire committee. Good Practice Notes are highlighted throughout and provide practical guidance to facilitate implementation. These were also developed through consensus of the entire committee.

<table>
<thead>
<tr>
<th>Recommendation category</th>
<th>Description</th>
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<tbody>
<tr>
<td>Evidence-based</td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>Body of evidence can be trusted to guide practice</td>
</tr>
<tr>
<td>B</td>
<td>Body of evidence can be trusted to guide practice in most situations</td>
</tr>
<tr>
<td>C</td>
<td>Body of evidence provides some support for recommendation(s) but care should be taken in its application</td>
</tr>
<tr>
<td>D</td>
<td>The body of evidence is weak and the recommendation must be applied with caution</td>
</tr>
<tr>
<td>Consensus-based</td>
<td>Recommendation based on clinical opinion and expertise as insufficient evidence available</td>
</tr>
<tr>
<td>Good Practice Note</td>
<td>Practical advice and information based on clinical opinion and expertise</td>
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</tbody>
</table>
Appendix D Full Disclaimer

This information is intended to provide general advice to practitioners, and should not be relied on as a substitute for proper assessment with respect to the particular circumstances of each case and the needs of any patient.

This information has been prepared having regard to general circumstances. It is the responsibility of each practitioner to have regard to the particular circumstances of each case. Clinical management should be responsive to the needs of the individual patient and the particular circumstances of each case.

This information has been prepared having regard to the information available at the time of its preparation, and each practitioner should have regard to relevant information, research or material which may have been published or become available subsequently.

Whilst the College endeavours to ensure that information is accurate and current at the time of preparation, it takes no responsibility for matters arising from changed circumstances or information or material that may have become subsequently available.