Abstract:

SOMANZ (Society of Obstetric Medicine Australia and New Zealand) has written a guideline to provide evidence based guidance for the investigation and care of women with sepsis in pregnancy or the postpartum period. The guideline is evidence based and incorporates recent changes in the definition of sepsis. The etiology, investigation and treatment of bacterial, viral and non-infective causes of sepsis are discussed. Obstetric considerations relevant to anesthetic and intensive care treatment in sepsis are also addressed. A multi-disciplinary group of clinicians with experience in all aspects of the care of pregnant women have contributed to the development of the guidelines. This is an executive summary of the guidelines.

Background

Despite an overall decline in maternal mortality in Australia, the maternal mortality rate from sepsis has increased from 0.6 per 100,000 in 2003-2005 to 0.8 per 100,000 in 2008-2012. In the period 2008-2012, sepsis accounted for 11.4% of maternal deaths in Australia. Group A beta hemolytic streptococcal (GAS) infection is the most common pathogen resulting in 25% of maternal deaths from sepsis in both Australia and the UK. The ninth Perinatal and Maternal Mortality Review Committee: report on maternal deaths in NZ between 2006-2013 indicates 50% of deaths from sepsis were related to Group A streptococcus. Sepsis continues to be one of the major causes of maternal mortality among Aboriginal and Torres Strait Islander women.

In Australia, the Clinical Excellence Commission developed the “Sepsis Kills” program to reduce poor outcomes from sepsis by improving recognition and management. A specific maternal pathway was created. Sepsis can arise at any time during pregnancy and is often associated with a delay in diagnosis. The normal physiological changes of pregnancy may mask early signs of sepsis. The woman has a unique ‘organ perfusion monitor’, namely the
fetus. Maternal sepsis with or without hemodynamic instability may present with fetal distress as the uteroplacental circulation is not auto-regulated. Thus any maternal circulatory insufficiency arising from sepsis may result in compromised fetal perfusion. Management plans need to take into consideration the altered immunological response of the woman and altered physiological responses during pregnancy. Consideration needs to be also given to the impact of the condition as well as the effect of its treatment on the fetus. The additional complexity of sepsis therapy in breast-feeding women also requires appropriate consideration.

These are the recommendations of a multidisciplinary working party convened by the Society of Obstetric Medicine of Australia and New Zealand (SOMANZ).

Methods

The clinical evidence for this guideline has been analysed by the authors according to the GRADE (Grading of Recommendations Assessment, Development and Evaluation) system. The principles followed in the development of these guidelines include: 1) a systematic review of the evidence with at least the following searches deployed: Cochrane library, Medline and EMBASE. 2) The authors formed a multi-disciplinary team 3) Financial support for administration was provided by SOMANZ. None of the authors received any direct financial compensation for their time to the development of this guideline. Where there were conflicting views, three authors (HR, LB and AM) reviewed the topic and came to a consensus. All authors reviewed the final guideline.

Definition of sepsis in pregnancy

Despite significant advances, understanding of the pathobiology of sepsis remains incomplete and currently no gold standard diagnostic test exists to confirm the presence of sepsis. Sepsis is broadly defined as life-threatening organ dysfunction caused by a dysregulated host response to infection. It is this dysregulated response and subsequent organ dysfunction that
differentiates sepsis from infection. The clinical signs may be insidious until sepsis is far advanced, which may occur very rapidly. Therefore early detection of sepsis is essential for appropriate multidisciplinary management to ensure the best outcomes for the mother and her baby. Septic patients may progress to develop septic shock, multi organ failure and death.

Septic shock is defined as a subset of sepsis in which profound circulatory, cellular, and metabolic abnormalities substantially increase mortality. More recently, the Sequential (sepsis-related) Organ Failure Assessment score (SOFA) has been shown to be useful in identifying those patients with a suspected infection who are likely to have a prolonged ICU (Intensive Care Unit) stay or die in hospital. A q(quick) SOFA score can be used to identify patients promptly at the bedside using only clinical information.

**SOMANZ definition of sepsis**

Recognising the patient with sepsis is paramount and is the first step in appropriate assessment and management. The flowchart in figure 1 summarises the clinical steps involved in the assessment and management of sepsis in pregnancy.

We recommend using the obstetrically modified qSOFA (omqSOFA) (Table 1). The obstetrically modified SOFA (omSOFA) score should be used for a subsequent, more thorough, assessment.

**Obstetrically modified quick SOFA (omqSOFA) score**

The qSOFA requires only clinical data for assessment and thus can be performed quickly without waiting for the results of biochemical or laboratory tests. In the non-pregnant patient, this score incorporates: systolic blood pressure of 100mmHg or less, respiratory rate of 22/min or greater and altered mentation; Glasgow coma score (GCS) less than 15. For each variable present, a score of 1 is attributed resulting in a score range of 0 - 3. A qSOFA score of greater than or equal to 2 has predictive validity for discriminating patients with an increased risk of in-hospital mortality. However, the data from which the score was derived

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However, is retrospective and validated in a heterogeneous population, with an average age of 61 years, of whom half were male. Extrapolation to pregnant and postpartum women should be undertaken with caution.

A woman’s gravid state will impact some of the variables in the qSOFA. Pregnancy significantly affects systolic blood pressure but not respiratory rate \( \text{or mentation}. \) During pregnancy, although women normally develop a respiratory alkalosis and compensatory metabolic acidosis, this is not through an increase in respiratory rate. The higher respiratory rate of 25 breaths/min, as compared to 22/min derived from published literature, was agreed upon by the writing committee as it aligned with cut offs employed on the maternity observation charts. Respiratory rate is poorly assessed in routine clinical practice. Systolic blood pressure usually decreases by 5-10mmHg in pregnancy. Furthermore a significant number (approximately 15%) of the obstetric population will have a usual systolic blood pressure at any given gestation, of less than 100mmHg and normal pregnancy outcomes.

Given these changes, several modifications are suggested to the qSOFA criteria when applied to pregnancy; obstetrically modified qSOFA (omqSOFA). Using the omqSOFA (Table 1), sepsis (as distinct from infection) in pregnant women should be considered where 2 or more of the following are present:

- Systolic blood pressure of 90mmHg or less
- Respiratory rate of 25/min or greater
- Altered mentation (any state other than ‘Alert’ on maternal observation charts). GCS (Glasgow coma scores) are not typically formally assessed as part of routine observations in obstetric wards.

**Obstetrically modified SOFA (omSOFA) score**

If sepsis is suspected based on screening, then assessment for end-organ dysfunction should be undertaken. Sepsis has been defined as an acute change in the total SOFA score of \( \geq 2 \) points consequent to infection. The baseline SOFA score is generally assumed to be zero where there is no pre-existing organ dysfunction. In the general population a SOFA of \( \geq 2 \) is associated with an overall mortality of 10%, these data are yet to be validated in the obstetric population. The SOFA score, in observational data, has been found to be significantly higher.
in pregnant women who have died compared to those who have survived in an intensive care setting. However the SOFA score has not undergone appropriate prospective validation in pregnant and postpartum populations. The studies that exist to date have been small, retrospective and undertaken in resource challenged environments which may limit their applicability generally.

This guideline recommends several modifications when applying the SOFA score to pregnancy (obstetrically modified SOFA- omSOFA) as indicated (Table 2):

- In order to demonstrate evidence of end organ dysfunction a score of \( \geq 2 \) needs to be attained. Therefore, scores of 3 or 4 in each category have been removed for the purposes of simplification.
- During pregnancy, serum creatinine levels are significantly reduced with the normal range being 35-80 µmol/L. For practical purposes, the serum creatinine cut off for the scores of 0, 1 or 2 have been adjusted to <90 µmol/L, 90-120 µmol/L or greater than 120 µmol/L respectively.
- As the GCS is not routinely assessed on maternity wards, the central nervous system category has been changed to reflect the maternal observation charts in routine clinical use. Alert will be scored 0, rousable by voice as 1 and rousable only by pain as 2. Any score other than 0 or ‘alert’ should trigger a GCS to be performed.
- Healthy pregnant women may have a mean arterial pressure less than 70 mmHg. Thus the SOFA score should be interpreted in the context of the woman’s premorbid blood pressure.

**SOMANZ Definition of septic shock**

If sepsis progresses, septic shock may ensue and this is associated with a substantial increase in mortality compared to sepsis alone. The clinical criteria validated to identify septic shock in non-pregnant patients include:

- Hypotension requiring vasopressor therapy to maintain a MAP 65 mmHg or greater (despite adequate fluid resuscitation) and
- Serum lactate greater than 2 mmol/L after adequate fluid resuscitation

No alterations have been made to these definitions for pregnancy.
C-reactive protein (CRP) has not been included in the current definition. It lacks sensitivity and is highly variable in pregnancy and should not be used to define or diagnose sepsis \(^{21}\). Fever, although helpful in identifying pregnant patients with suspected infections or sepsis, has also not been included, as it lacks sensitivity\(^{25}\) (see section on fever below).

Given that maternal physiology gradually returns to normal postpartum, we recommend the definition of postpartum sepsis be the same as for non-pregnant patients after the first week postpartum.

### Key Points 1:
- Screen for sepsis using the omqSOFA score.
- Assess for evidence of end organ dysfunction using omSOFA score
- Septic shock is a complication of sepsis evidenced by the need for vasopressors and elevated lactate

**GRADE: MODERATE QUALITY EVIDENCE**

the health impacts on the offspring, Dreier and colleagues\(^{26}\) reported an increase in the rate of pregnancies affected by a neural tube defect in mothers who experienced fever in the first trimester or peri-conceptually (pooled odds ratio of 2.9 (95% Confidence Interval [CI] 2.22-3.79)). Oral clefts were also more common in mothers with fever early in pregnancy (OR 1.94, 95% CI 1.35-2.79). Congenital heart defects were weakly associated with early maternal fever early in pregnancy (OR 1.54, 95% CI 1.37-1.74). No effect of maternal fever was seen on the risk of miscarriage, stillbirth or preterm labour.

Human studies appear to vary from animal studies in the absence of a dose-response relationship. In animals where a temperature elevation of greater than 2°C was documented there was a greater risk of teratogenic outcome\(^{27}\). In human studies, reliance on self-reported temperature may lead to inaccurate results\(^{26}\).

Few studies are available to document the long term developmental outcomes of infants exposed to fever in pregnancy. In the CHARGE (Childhood Autism Risks from Genetics and Environment) study \(^{28}\), mothers of children diagnosed with autism spectrum disorder (ASD)
or developmental delay (DD) were asked retrospectively, 2 – 5 years after their pregnancy, whether or not they had suffered from flu or a fever in pregnancy. Neither ASD nor DD were associated with self-reported influenza in pregnancy. However, both ASD and DD were associated with the report of fever during pregnancy, OR 2.12 (95% CI 1.17-3.84) and OR 2.50 (95% CI 1.20 – 5.20) respectively. The offspring of mothers who took anti-pyretic medications had a lower risk of ASD, OR 1.30 (0.59-2.84) compared with those who did not OR 2.55 (1.30-4.99).

High-dose aspirin and non-steroidal anti-inflammatory agents should be used with caution during the third trimester due to the risk of premature closure of the fetal ductus arteriosus. Alternative agents should be considered.

Key Points 2:

- Fever may be associated with an increased risk of congenital abnormalities
- Antipyretics such as paracetamol, may be beneficial in reducing adverse pregnancy outcomes

GRADE: LOW QUALITY EVIDENCE

Although most commonly bacterial in etiology, sepsis can also result from viral and other causes (Table 3). A number of non-infective conditions can mimic sepsis and should be considered.

Key Points 3:

- Sepsis-related maternal death is most commonly caused by Group A Streptococcus
- E. Coli is the commonest cause of maternal bacterial infection
- Consider non-bacterial and non-infective conditions that may mimic sepsis

GRADE: HIGH QUALITY EVIDENCE
Investigations in sepsis

Obstetric patients with infections may present with nonspecific symptoms and early investigation is necessary to exclude severe infection. Maternity units should ensure observations are recorded on maternity specific charts for all obstetric and postpartum patients. Utilization of these charts has been shown to promote earlier detection and therefore treatment of pregnant or early post partum women who are developing a critical illness.

Once infection is suspected, treatment should be commenced and the likely source and severity of sepsis elucidated via history, examination and investigations. Investigations are directed at determining the etiology and risk stratification of sepsis as well as organisation of care for women in the most appropriate clinical area. Sepsis during pregnancy will require maternal investigations as detailed below and fetal wellbeing assessment with cardiotocograph (CTG) and/or ultrasound. Table 5 lists the first line investigations to be undertaken in sepsis, possible changes in sepsis and normal pregnancy reference ranges if different from the non pregnant state. Two important consideraions are:

- Two sets of peripheral blood cultures should be taken immediately and sequentially if possible before administration of antibiotics. Their collection should not delay antibiotic treatment. Cultures should also be taken from all vascular catheter lines (catheter blood cultures) and any other potential sources of infection. Determining the etiology of the infection will allow for targeting of antimicrobial therapy.
- Arterial blood gases should be undertaken to determine presence of hypoxia, hypercapnea, metabolic state and lactate level. Elevated lactate levels are an indication of tissue hypoperfusion with values greater than 2 mmol/L being associated with increased mortality in pregnancy. The sampling site does not affect the serum lactate results (eg. arterial, venous or capillary). If unable to obtain an arterial lactate, a venous lactate can be collected.

Additional investigations may be required depending on initial investigation results and the likely source of infection. Appropriate imaging should not be withheld because a woman is pregnant. Concerns about radiation exposure to her developing fetus or proliferating maternal breast tissue need to be balanced against the clinical need for imaging.

Key Points 4:

- Maternal observation should be recorded on maternity specific observation charts
- Blood cultures and other specimens should be obtained before antibiotics given:
Treatment in the golden hour

Treatment should be commenced as soon as practical - ideally within the first hour (‘golden hour’) of sepsis being suspected. Empiric treatment should include fluid resuscitation, correction of hypoxia and antimicrobials (antibiotics or antivirals as appropriate). Source control, consideration of the appropriate area of care as well as venous thromboembolism prophylaxis also need to be considered.

Fluid resuscitation

In most circumstances, the initial treatment of hypotension in sepsis is fluid administration. Fluid resuscitation is vital in sepsis to optimise circulating volume and improve blood pressure and tissue perfusion. The preferred fluid for resuscitation is an isotonic crystalloid (usually 0.9% normal saline). Blood may also be used as fluid replacement if there is evidence of blood loss or severe anaemia. If, despite fluid resuscitation (1-2L), the mean arterial pressure and other indices do not improve, care will need to be escalated. Vasopressors may be required see ‘Intensive Care Issues’ below.

Treatment for bacterial sepsis

When a bacterial source of sepsis in pregnancy or postpartum is suspected, prompt treatment with antibiotics within one hour is critical. Maternal mortality can increase by 8% for each hour’s delay in administering antibiotics. Following principles of antibiotic stewardship recommended in both Australia and New Zealand, selection of antibiotics needs to be
appropriate for the suspected infection, whilst minimizing the risk of adverse effects and reducing the emergence of antibiotic resistance.\textsuperscript{37, 38}

A consultant obstetrician and a physician experienced in the management of sepsis in pregnancy should be involved in the care of a pregnant or postpartum patient from the time of sepsis diagnosis or recognition. However, investigation and treatment should not be delayed while waiting for expert consultation. Once a source of sepsis is identified, source control is a priority and may involve abscess drainage or delivery of the fetus.

When treating immunosuppressed women a physician should be involved in decision making as soon as possible after the diagnosis or recognition of sepsis. This should not delay antibiotic treatment. Women may be immunosuppressed with immunomodulators for solid organ transplant, malignancy or autoimmune disease; chronic infection including HIV; or significant medical co-morbidities such as diabetes.

**Treatment of sepsis of unknown source**

Table 6 summarizes the recommendations for empiric antibiotic treatment. Antibiotic resistance patterns and local practices will vary so it is important to seek local specialist advice as soon as possible. Therapy should be refined as culture and imaging results define the source of sepsis and its culprit organisms. The Australia-wide eTG (electronic antibiotic Therapeutic Guidelines) are available at the following URL: https://tgldedp.tg.org.au/guideLine?guidelinePage=Antibiotic&frompage=etgcomplete

**Treatment of sepsis of viral etiology – Influenza**

Neuraminidase inhibitors are recommended for the treatment of influenza.\textsuperscript{39} Oseltamivir (treatment dose: 75mg capsule twice daily for 5 days) has more obstetric safety data available than zanamivir and is the agent of choice in pregnancy.\textsuperscript{40} There is a low rate of transplacental transfer, estimated at between 1-14\% of maternal concentrations in ex vivo perfusion studies.\textsuperscript{41} In the setting of H1N1 pandemic influenza, early antiviral therapy (initiation < 2 days) in pregnant women was associated with an 84\% reduction in admissions to intensive care.\textsuperscript{42}
Thromboembolism prophylaxis

Both pregnancy and sepsis are independent risk factors for venous thromboembolism\textsuperscript{43}. Thus prevention of venous thromboembolism is critically important. Unfractionated heparin (UH) and low molecular weight heparin (LMWH) have been used extensively in pregnancy and have been shown in large clinical trials to be effective in the prevention of thromboembolism\textsuperscript{44}.

Pharmacological considerations for pregnancy and breastfeeding

While most antibiotics are found in breast milk of a lactating woman, the relative infant dose is generally small. Breastfed infants should be monitored for side effects such as diarrhoea, vomiting, skin rash or thrush while their mothers are being treated with antibiotics. Please refer to the eTG for advice on antibiotic use in breast feeding mothers, URL: https://tgldedp.tg.org.au.acs.hcn.com.au/quicklinks?type=Pregnancyandbreastfeeding.

Timing and mode of delivery

The timing of delivery will be determined by a) the presence of intrauterine sepsis, b) the nature of the maternal sepsis and response to initial resuscitation efforts and c) the gestation of the pregnancy and fetal status.

In the setting of intrauterine sepsis, delivery should always be considered regardless of the gestation. Corticosteroids should be considered for fetal lung maturation but this decision needs to be balanced against the urgency of delivery\textsuperscript{45}.

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Intrauterine sepsis should be suspected in the presence of maternal fever, ruptured membranes or recent intrauterine procedures such as amniocentesis, maternal tachycardia, fetal tachycardia, uterine tenderness or offensive vaginal discharge. While the onset of chorioamnionitis may be non-specific and insidious, rapid deterioration is possible.

In cases of extra-uterine sepsis, efforts to treat maternal sepsis and prolong gestation should be considered at gestations remote from term, although it is reasonable to consider delivery in term pregnancies as a means of improving maternal resuscitation efforts. Fetal wellbeing should be monitored during maternal sepsis with the most gestation appropriate method.

**Role of the anesthetist in managing maternal sepsis**

The role of the anesthetist in managing maternal sepsis includes:

- initial resuscitation and stabilisation of the patient
- transfer of the sick patient (to imaging or intensive care unit)
- intra-operative care during delivery
- anaesthesia for surgical management of sepsis

**Patient transfer**

The transport team must be experienced in securing airways, ventilation, resuscitation and other anticipated emergency procedures. A comprehensive hand over should take place at the receiving facility. The responsibility of intra and inter-hospital transfers often rests with the anesthetist to provide the standard of care as set out in The Australian and New Zealand College of Anesthetists professional document, PS52 (2015) Guidelines for Transport of
Critically Ill Patients. These guidelines state, “the level of care provided during transport must aim to at least equal that at the point of referral and must prepare the patient for admission to the receiving services” 47.

**Spinal anaesthesia**

Underlying sepsis is a risk factor for infectious complications of regional anaesthesia. The greatest concern is infection around the spine and spinal cord, presenting either as meningitis or cord compression secondary to abscess formation with the potential for permanent neurological deficit. It is difficult to accurately quantify the risk of infection following neuraxial analgesia and anaesthesia due to the great variability in the reported incidence in the literature; from an estimated incidence of spinal/epidural abscess after epidural analgesia of 1 in 1,930 to 1.1 infections per 100,000 neuraxial blocks 48. However, epidural anaesthesia carries a greater risk of infectious complications than spinal techniques 49.

Despite the low risk of central nervous system infection, which may potentially occur in any bacteraemic patient, the decision to proceed with neuraxial blockade in a febrile or infected patient must be carefully considered and made on a case by case basis 50-52:

1. Except in the most extraordinary circumstances, central neuronal block should not be performed in patients with untreated systemic infection.
2. Patients with evidence of systemic infection may safely undergo spinal anaesthesia, provided appropriate antibiotic therapy is initiated before dural puncture and the patient has shown a response to therapy; placement of an indwelling epidural or intrathecal catheter remains controversial.
3. Spinal anaesthesia may be safely performed in patients at risk for low grade transient bacteraemia after dural puncture.

**General anaesthesia**

The septic obstetric patient often exhibits hemodynamic instability and has a greater (than just pregnancy induced) metabolic oxygen demand. In practice, relevant issues for the administration of general anaesthesia include:

- **A – Airway**: Delayed gastric emptying with increased risk of reflux and aspiration. It is recommended that women are premedicated with combination antacid antihistamine prophylaxis, e.g. effervescent ranitidine 150mg. A rapid sequence induction is also recommended.
• **B – Breathing**: Increased metabolic demand leading to accelerated hypoxaemia during periods of apnoea. It is recommended women receive adequate pre-oxygenation prior to anaesthesia induction. There is reduced functional residual capacity with increased ventilation/perfusion mismatch. It is recommended ventilation strategies to maintain oxygenation and minimise further lung injury are employed.

• **C – Circulation**: Maintenance of systemic blood pressure for adequate organ perfusion, including the placental bed. It is recommended to avoid aortocaval compression using a lateral uterine tilt, ensure adequate fluid resuscitation including the appropriate use of blood products and if required, inotropic support. Alpha adrenergic agonists (specifically noradrenaline) are the agents of choice for maintenance of uteroplacental flow.

### Intensive Care Issues

Adequate initial resuscitation and treatment of sepsis may result in stabilisation and prevent progression and deterioration, averting the requirement for intensive care. However, as stated early consideration should be given to establishing the most appropriate venue of care for these women. The indications for ICU admission will vary depending on local resources and expertise. It is preferable that an ICU opinion and/or admission should occur before the development of severe complications such as frank organ failure or catastrophic shock. If any clinical concern exists then escalation of care should be considered early. Table 7 sets out in general terms when an ICU opinion should be obtained and an ICU admission considered.

### Key Points 7:
- Neuraxial blocks in women with untreated sepsis are associated with increased complications
- Septic women experience increased haemodynamic instability during anesthesia

GRADE: LOW QUALITY EVIDENCE

### Key Points 8:
- Liaise with ICU where there is evidence of organ dysfunction, hypotension or cardio respiratory compromise
- Early ICU involvement is preferable and encouraged
- Indications for ICU admission varies based on local resources and experience

GRADE: LOW QUALITY EVIDENCE
Table and Figure Legends

**Figure 1:** Flowchart for the assessment and management of sepsis in pregnancy. The clinically important steps are noted in blue boxes and comments relating to the steps are in neighboring green boxes.

**Table 1:** Obstetrically modified qSOFA score
**Table 2:** Obstetrically modified SOFA score
**Table 3:** Infectious causes of sepsis in pregnancy and postpartum
**Table 4:** Non-infectious conditions that can mimic sepsis in pregnancy
**Table 5:** First line investigations recommended for suspected sepsis
**Table 6:** Recommendations for antimicrobial treatment of sepsis with unknown source
**Table 7:** Indications for involvement of ICU
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Table 1: Obstetrically modified qSOFA score

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
</tr>
<tr>
<td><strong>Systolic Blood Pressure</strong></td>
<td>≥90mmHg</td>
</tr>
<tr>
<td><strong>Respiratory Rate</strong></td>
<td>Less than 25 breaths/min</td>
</tr>
<tr>
<td><strong>Altered mentation</strong></td>
<td>Alert</td>
</tr>
</tbody>
</table>

mmHg- Millimetres of mercury, Min- minute
Table 2: Obstetrically modified SOFA score

<table>
<thead>
<tr>
<th>System Parameter</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
</tr>
<tr>
<td><strong>Respiration</strong></td>
<td></td>
</tr>
<tr>
<td>PaO₂/FIO₂</td>
<td>≥400</td>
</tr>
<tr>
<td><strong>Coagulation</strong></td>
<td></td>
</tr>
<tr>
<td>Platelets x10⁶/L</td>
<td>≥150</td>
</tr>
<tr>
<td><strong>Liver</strong></td>
<td></td>
</tr>
<tr>
<td>Bilirubin (μmol/L)</td>
<td>≤20</td>
</tr>
<tr>
<td><strong>Cardiovascular</strong></td>
<td></td>
</tr>
<tr>
<td>Mean Arterial Pressure (mm Hg)</td>
<td>MAP≥70</td>
</tr>
<tr>
<td><strong>Central Nervous System</strong></td>
<td>Alert</td>
</tr>
<tr>
<td><strong>Renal</strong></td>
<td></td>
</tr>
<tr>
<td>Creatinine (μmol/L)</td>
<td>≤90</td>
</tr>
</tbody>
</table>

MAP- Mean arterial pressure, mmHg – millimetres of mercury, μmol/L- micromoles per litre, PaO₂- partial pressure of oxygen (in mmHg), FIO₂- fraction of inspired oxygen (expressed as a decimal)
Table 3: Infectious causes of sepsis in pregnancy and postpartum

<table>
<thead>
<tr>
<th>Infection</th>
<th>Pathogens</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bacterial - common</strong></td>
<td>Group A- beta-hemolytic Streptococcus (GAS) pyogenes</td>
</tr>
<tr>
<td></td>
<td>Escherichia Coli</td>
</tr>
<tr>
<td></td>
<td>Group B Streptococcus</td>
</tr>
<tr>
<td></td>
<td>Klebsiella pneumoniae</td>
</tr>
<tr>
<td></td>
<td>Staphylococcus aureus</td>
</tr>
<tr>
<td></td>
<td>Streptococcus pneumonia</td>
</tr>
<tr>
<td></td>
<td>Proteus mirabilis</td>
</tr>
<tr>
<td></td>
<td>Anaerobic organisms</td>
</tr>
<tr>
<td><strong>Bacterial – less common</strong></td>
<td>Haemophilus influenza</td>
</tr>
<tr>
<td></td>
<td>Listeria monocytygenes</td>
</tr>
<tr>
<td></td>
<td>Clostridium species</td>
</tr>
<tr>
<td></td>
<td>MycobacteriumTuberculosis</td>
</tr>
<tr>
<td><strong>Viral</strong></td>
<td>Influenza</td>
</tr>
<tr>
<td></td>
<td>Varicella zoster virus</td>
</tr>
<tr>
<td></td>
<td>Herpes Simplex virus</td>
</tr>
<tr>
<td></td>
<td>Cytomegalovirus</td>
</tr>
</tbody>
</table>
Table 4: Non-infectious conditions that can mimic sepsis in pregnancy

<table>
<thead>
<tr>
<th>Condition</th>
<th>Common Maternal Clinical Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute pulmonary embolism</td>
<td>Hypotension, tachypnoea, tachycardia, low grade fever</td>
</tr>
<tr>
<td>Amniotic fluid embolism</td>
<td>Hypotension, tachycardia, haemorrhage</td>
</tr>
<tr>
<td>Acute pancreatitis</td>
<td>Fever, nausea, vomiting, abdominal pain</td>
</tr>
<tr>
<td>Acute Fatty Liver of Pregnancy</td>
<td>Fatigue, nausea, vomiting, abdominal pain, jaundice, impaired level of consciousness</td>
</tr>
<tr>
<td>Adverse drug reactions, drug fever</td>
<td>Hypotension, relative bradycardia, fever, rash, angio-oedema</td>
</tr>
<tr>
<td>Acute liver failure-drug related, viral</td>
<td>Jaundice, nausea, vomiting, abdominal pain, impaired level of consciousness</td>
</tr>
<tr>
<td>Acute adrenal insufficiency</td>
<td>Weakness, fatigue, nausea, anorexia, weight loss, hypotension, fever</td>
</tr>
<tr>
<td>Acute pituitary insufficiency</td>
<td>Failure to lactate, hypotension, relative bradycardia, polyuria, polydipsia</td>
</tr>
<tr>
<td>Autoimmune conditions</td>
<td>Low grade fever, rash (eg. malar rash), arthritis, dry eyes or mouth, mouth ulcers, diagnostic serology</td>
</tr>
<tr>
<td>Concealed haemorrhage including ectopic pregnancy</td>
<td>Hypotension, tachycardia, low grade fever</td>
</tr>
<tr>
<td>Disseminated Malignancy</td>
<td>Low grade fever, weight loss</td>
</tr>
<tr>
<td>Pelvic Thrombosis</td>
<td>Pelvic pain, fever,</td>
</tr>
<tr>
<td>Transfusion reactions</td>
<td>High fever, rigors, dysrhythmia, tachypnoea, hypotension, rash, bleeding, haematuria</td>
</tr>
</tbody>
</table>
### Table 5: First line investigations recommended for suspected sepsis

<table>
<thead>
<tr>
<th>Investigation</th>
<th>Obstetric reference range (if relevant)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blood cultures</strong></td>
<td></td>
</tr>
<tr>
<td>- At least 2 sets, prior to antibiotic commencement as long as there is no delay.</td>
<td></td>
</tr>
<tr>
<td>- Obtain samples from different sites</td>
<td></td>
</tr>
<tr>
<td>- Cultures should also be obtained from IV access devices</td>
<td></td>
</tr>
<tr>
<td><strong>Other Cultures</strong></td>
<td></td>
</tr>
<tr>
<td>- Obtain cultures of additional sites as clinically indicated and as soon as possible</td>
<td></td>
</tr>
<tr>
<td>Eg. urine MCS, wound swab - episiotomy, caesarean, placental swabs, amniotic fluid, sputum culture, naso-pharyngeal aspirate/swab, cerebrospinal fluid, vaginal swabs, stool culture</td>
<td></td>
</tr>
<tr>
<td><strong>Arterial blood gases</strong></td>
<td></td>
</tr>
<tr>
<td>- detect acidosis, hypoxaemia, lactate as below</td>
<td></td>
</tr>
<tr>
<td><strong>Lactate</strong></td>
<td></td>
</tr>
<tr>
<td>- elevated levels in sepsis relate to tissue hypoperfusion and are associated with an increased sepsis mortality risk</td>
<td></td>
</tr>
<tr>
<td><strong>Full blood count</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Coagulation studies</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Creatinine, urea and electrolytes</strong></td>
<td></td>
</tr>
<tr>
<td>- Measure at baseline and until the patient improves, elevated creatinine is a sign of severe sepsis</td>
<td></td>
</tr>
<tr>
<td><strong>Liver function tests</strong></td>
<td></td>
</tr>
<tr>
<td>- Baseline test, may be elevated if sepsis source is from hepatic or perihepatic infections</td>
<td></td>
</tr>
<tr>
<td>- May be elevated due to septic shock affecting hepatic blood flow and metabolism</td>
<td></td>
</tr>
</tbody>
</table>

PaO<sub>2</sub>: 1<sup>st</sup> trimester: 93-100 mmHg, 2<sup>nd</sup> trimester: 90-98 mmHg, 3<sup>rd</sup> trimester: 92-107 mmHg
PaCO<sub>2</sub>: 25-33 mmHg, Arterial pH: 7.4-7.47, HCO<sub>3</sub>: 16-22 mmol/L, 0.6-1.8 mmol/L

White cell count: 6-17 × 10<sup>9</sup>/L (may increase to 9-15 × 10<sup>9</sup>/L immediately post-delivery). Steroids also increase white cell count
Platelets – lower limit of normal 150-420 ×10<sup>9</sup>/L

No change

Creatinine Varies with Gestation (reference ranges): 1<sup>st</sup> trimester: 35-62 μmol/L, 2<sup>nd</sup> trimester: 35-71 μmol/L, 3<sup>rd</sup> trimester: 35-80 μmol/L
AST 3-33 U/L, ALT 2-33 U/L, Alkaline Phosphatase 17-229 U/L
GGT 2-26 U/L, Total Bilirubin 1.7-19 μmol/L
CXR

Fetal Assessment – CTG and/or fetal ultrasound

A non-reassuring CTG suggests inadequate uteroplacental perfusion and may reflect maternal organ hypoperfusion or intrauterine sepsis

\( \text{PaO}_2 \) - partial pressure of oxygen (in mmHg), \( \text{PaCO}_2 \) - partial pressure of carbon dioxide (in mmHg), \( \text{HCO}_3^- \) - Bicarbonate, IV - intravenous, MCS - microscopy, culture and sensitivity, CTG - cardiotocograph, CXR - Chest x-ray, mmol/L - millimoles per litre, \( \mu \text{mol/L} \) - micromoles per litre, AST - aspartate aminotransferase, ALT - alanine aminotransferase, GGT - gamma glutamyl transferase, U/L - units per litre.
<table>
<thead>
<tr>
<th>Community-acquired sepsis (source not apparent)</th>
<th>Australian and New Zealand Antibiotic Regimen⁹</th>
<th>Alternative for penicillin hypersensitivity⁹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aus : amoxicillin/ampicillin 2g IV 6-hourly PLUS gentamicin 4.7mg/kg (first dose) IV* PLUS metronidazole 500mg IV 12-hourly</td>
<td>Clindamycin 600mg IV 8-hourly PLUS gentamicin 4.7mg/kg (first dose) IV (severe hypersensitivity)</td>
<td>Cefazolin 2g IV 6-hourly PLUS gentamicin 4.7mg/kg (first dose) IV* PLUS metronidazole 500mg IV 12-hourly (mild-moderate hypersensitivity)</td>
</tr>
<tr>
<td>NZ: cefuroxime 1.5g IV 8-hourly PLUS gentamicin 4.7mg/kg (first dose) IV* PLUS metronidazole 500mg IV 12-hourly</td>
<td></td>
<td></td>
</tr>
<tr>
<td>At risk of MRSA sepsis (based on previous swabs/cultures and local epidemiology): ADD vancomycin 25-30mg/kg (loading dose) IV</td>
<td></td>
<td></td>
</tr>
<tr>
<td>At risk of Group A Streptococcal (GAS) sepsis: ADD clindamycin 600mg IV 8-hourly, PLUS consider normal immunoglobulin 1-2g/kg IV, for up to 2 doses during the first 72 hours</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Hospital-acquired sepsis (source not apparent)</th>
<th>Australian and New Zealand Antibiotic Regimen⁹</th>
<th>Alternative for penicillin hypersensitivity⁹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aus: piperacillin 4g + tazobactam 0.5g IV 8-hourly AND consider gentamicin 4.7mg/kg (first dose) IV* (if local epidemiology suggests Gram negative aminoglycoside susceptibility)</td>
<td>Severe: ciprofloxacin 400mg IV 8-hourly PLUS vancomycin 25-30mg/kg IV*</td>
<td></td>
</tr>
<tr>
<td>NZ: cefuroxime 1.5g IV 8-hourly PLUS gentamicin 4.7mg/kg (first dose) IV* PLUS metronidazole 500mg IV 12-hourly</td>
<td></td>
<td></td>
</tr>
<tr>
<td>At risk of MRSA sepsis (based on previous swabs/cultures and local epidemiology or if line sepsis) ADD vancomycin 25-30mg/kg (loading dose) IV</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th><strong>At risk of multidrug-resistant Gram-negative organisms</strong>: use as a SINGLE AGENT</th>
<th><strong>meropenem</strong> 1g IV 8-hourly</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>At risk of Group A Streptococcal (GAS) sepsis</strong>: ADD <strong>clindamycin</strong></td>
<td>600mg IV 8-hourly PLUS consider normal immunoglobulin 1-2g/kg IV, for up to 2 doses during the first 72 hours</td>
</tr>
<tr>
<td><strong>Consider</strong> <strong>influenza</strong></td>
<td><strong>Oseltamivir</strong> 75mg BD or <strong>Zanamivir</strong> 2 inhalations (each 5mg) twice daily for 5 days</td>
</tr>
</tbody>
</table>

* Use local protocols for gentamicin and vancomycin dosing and monitoring. Once daily dosing of gentamicin in pregnancy and postpartum can be used and in pregnancy results in levels below the toxicity threshold for more hours per day than in 8-hourly dosing. *NZ regime does not cover listeria – if suspected use a penicillin as per Australian regime. *NZ has increasing Group B strep resistance to clindamycin and macrolides – if penicillin hypersensitivity seek expert advice for best agent.  

**IV-** intravenous, **MRSA-** methicillin resistant staphylococcus aureas, **mg/kg-** milligrams per kilogram, **BD-** twice daily, **mg-** milligrams.

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**Table 6: Recommendations for antimicrobial treatment of sepsis with unknown source**

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Table 7: Indications for involvement of ICU

<table>
<thead>
<tr>
<th>Indications for ICU involvement</th>
<th>Signs or observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiorespiratory compromise</td>
<td>hypotension, circulatory instability, worsening tachypnoea, worsening hypoxia, increasing supplemental oxygen requirements</td>
</tr>
<tr>
<td>Evidence of organ dysfunction</td>
<td>altered mental status, oliguria, worsening urea and creatinine, other e.g. coagulation failure, cytopenias, worsening hepatic dysfunction</td>
</tr>
<tr>
<td>Other evidence of hypoperfusion</td>
<td>metabolic/lactic acidosis, signs of poor tissue perfusion, signs of inadequate placental perfusion</td>
</tr>
<tr>
<td>Other serious clinical concern</td>
<td></td>
</tr>
</tbody>
</table>