Perinatal Practice Guideline

Clinical Guideline
Sepsis in Pregnancy

Policy developed by: SA Maternal, Neonatal & Gynaecology Community of Practice
Approved SA Health Safety & Quality Strategic Governance Committee on: 01 March 2017
Next review due: 31 March 2020

Summary
The purpose of this guideline is to provide clinicians with information and treatment guidelines for sepsis in pregnancy. It particularly focuses on recognition of risk factors, diagnosis and management of sepsis in the peri partum period, including Early Goal Directed Therapy in the case of severe sepsis.

Keywords
Sepsis in Pregnancy, sepsis, chorioamnionitis, infection, antibiotics, toxic shock syndrome, clinical guideline, PPG, perinatal practice guideline, early goal directed therapy, toxic shock syndrome, endometritis, bacteraemia, systemic inflammatory response syndrome, septicemia

Policy history
Is this a new policy? N
Does this policy amend or update an existing policy? Y v1.0
Does this policy replace an existing policy? Y
If so, which policies?

Applies to
All Health Networks
CALHN, SALHN, NALHN, CHSALHN, WCHN, SAAS

Staff impact
All Staff, Management, Students, All Clinical, Medical, Midwifery, Nursing, Allied Health, Emergency, Mental Health, Pathology

PDS reference
CG190

Version control and change history

<table>
<thead>
<tr>
<th>Version</th>
<th>Date from</th>
<th>Date to</th>
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<tr>
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South Australian Perinatal Practice Guidelines

Sepsis in Pregnancy

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Note

This guideline provides advice of a general nature. This statewide guideline has been prepared to promote and facilitate standardisation and consistency of practice, using a multidisciplinary approach. The guideline is based on a review of published evidence and expert opinion.

Information in this statewide guideline is current at the time of publication.

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Health practitioners in the South Australian public health sector are expected to review specific details of each patient and professionally assess the applicability of the relevant guideline to that clinical situation.

If for good clinical reasons, a decision is made to depart from the guideline, the responsible clinician must document in the patient’s medical record, the decision made, by whom, and detailed reasons for the departure from the guideline.

This statewide guideline does not address all the elements of clinical practice and assumes that the individual clinicians are responsible for discussing care with consumers in an environment that is culturally appropriate and which enables respectful confidential discussion. This includes:

- The use of interpreter services where necessary,
- Advising consumers of their choice and ensuring informed consent is obtained,
- Providing care within scope of practice, meeting all legislative requirements and maintaining standards of professional conduct, and
- Documenting all care in accordance with mandatory and local requirements

Explanation of the Aboriginal artwork:
The Aboriginal artwork used symbolises the connection to country and the circle shape shows the strong relationships amongst families and the Aboriginal culture. The horse shoe shape design shown in front of the generic statement symbolises a woman and those enclosing a smaller horse shoe shape depicts a pregnant woman. The smaller horse shoe shape in this instance represents the unborn child. The artwork shown before the specific statements within the document symbolises a footprint and demonstrates the need to move forward together in unison.

Australian Aboriginal Culture is the oldest living culture in the world yet Aboriginal people continue to experience the poorest health outcomes when compared to non-Aboriginal Australians. In South Australia, Aboriginal women are 2-5 times more likely to die in childbirth and their babies are 2-3 times more likely to be of low birth weight. The accumulative effects of stress, low socio economic status, exposure to violence, historical trauma, culturally unsafe and discriminatory health services and health systems are all major contributors to the disparities in Aboriginal maternal and birthing outcomes. Despite these unacceptable statistics the birth of an Aboriginal baby is a celebration of life and an important cultural event bringing family together in celebration, obligation and responsibility. The diversity between Aboriginal cultures, language and practices differ greatly and so it is imperative that perinatal services prepare to respectively manage Aboriginal protocol and provide a culturally positive health care experience for Aboriginal people to ensure the best maternal, neonatal and child health outcomes.

Purpose and Scope of PPG

The purpose of this guideline is to provide clinicians with information and treatment guidelines for sepsis in pregnancy. It particularly focuses on recognition of risk factors, diagnosis and management of sepsis in the peri partum period, including Early Goal Directed Therapy in the case of severe sepsis.
Flowchart: Initial management of maternal sepsis

Call for help
Senior midwives, senior obstetrician/s, intensivist and anaesthetist

Monitoring
Observations
Respiratory rate, blood pressure, heart rate, SpO2, temperature

Fetal monitoring (if appropriate)
Fetal heart rate and / or continuous CTG

Rapid detection and response chart
Initial observations every 15 minutes

Investigations
Full clinical examination
Head to toe examination

Microbiology
Blood cultures, midstream urine, high vaginal swabs, wound swab, throat swab, stool sample

Blood tests
Complete blood picture, urea and electrolytes, coagulation profile, lactate, arterial blood gas

Imaging
Chest Xray, abdominal and pelvic ultrasound and CT scan

Investigations

Treatment
Fluid resuscitation
At least 20 mL/kg crystalloid

Broad-spectrum IV antibiotics
Consult with microbiologist or infectious diseases consultant as required

Remove source
e.g. evacuation of retained products, drainage of pelvic collection

Plan for labour / birth
(if appropriate) involve neonatologist

Intensive care support

Urinary catheter
Hourly urine output

Invasive monitoring
Consider central venous line and arterial line

Fetal monitoring
Flowchart: Initial management of maternal sepsis

Call for help
Senior midwives, senior obstetrician/s, intensivist and anaesthetist

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Intensive care support

Urinary catheter
Hourly urine output

Invasive monitoring
Consider central venous line and arterial line
Flowchart: Early goal directed therapy for sepsis during pregnancy

Suspected infection

The high risk patient: SBP < 90 mm Hg after 20-40 mL / kg volume challenge or serum lactate > 4 mmol / L

Antibiotics within 1 hour and source control

CVP

< 8 mm Hg

Crystalloid

> 8-12 mm Hg

MAP

< 65 or > 90 mm Hg

Vasoactive agents

> 65-90 mm Hg

SvO₂

< 70%

Packed RBC to Hct > 30%

> 70%

Inotrope(s)

Goals achieved

Decrease oxygen consumption

No

Suspected infection

The high risk patient: SBP < 90 mm Hg after 20-40 mL / kg volume challenge or serum lactate > 4 mmol / L

Antibiotics within 1 hour and source control

CVP

< 8 mm Hg

Crystalloid

> 8-12 mm Hg

MAP

< 65 or > 90 mm Hg

Vasoactive agents

> 65-90 mm Hg

SvO₂

< 70%

Packed RBC to Hct > 30%

> 70%

Inotrope(s)

Goals achieved

Decrease oxygen consumption

No
EGDT (early goal directed therapy) - goals and normal values in pregnancy chart³

<table>
<thead>
<tr>
<th>Measures</th>
<th>Resuscitation goals</th>
<th>Normal third trimester physiologic values</th>
</tr>
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<tr>
<td>Central venous pressure</td>
<td>8-12 mm Hg</td>
<td>4-10 mm Hg</td>
</tr>
<tr>
<td>Mean arterial pressure</td>
<td>65 mm Hg or more</td>
<td>84-96 mm Hg</td>
</tr>
<tr>
<td>Urine output</td>
<td>More than 0.5 mL / kg / hour</td>
<td>Minimum 0.5 mL / kg / hour</td>
</tr>
<tr>
<td>Mixed venous oxygen saturation</td>
<td>More than 70 %</td>
<td>More than 80 %</td>
</tr>
<tr>
<td>Heart rate</td>
<td>Decreasing in response to treatment</td>
<td>83 (+/- 10) beats / minute</td>
</tr>
</tbody>
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References

Useful online references

Acknowledgements

Summary of Practice Recommendations

> Sepsis may arise from many sources and is not limited to infections arising from the genital tract.
> The onset of sepsis may be insidious; women with severe sepsis may appear very well before suddenly collapsing, with little or no warning.
> Abdominal pain, fever (> 38°C), tachypnoea and sustained tachycardia (> 90 beats per minute) are indications for admission and intravenous antibiotics.
> Diarrhoea and/or vomiting in pregnant women may be serious signs of sepsis and an indication for commencing antibiotic therapy.
> Early investigation of non-specific symptoms in pregnant women is necessary to exclude serious infection.
> Urinary tract infection and chorioamnionitis are common infections associated with septic shock in the pregnant woman.
> Signs of severe sepsis in peripartum women, particularly with confirmed or suspected Group A streptococcal infection, should be regarded as an obstetric emergency.
> Early involvement of senior obstetrician / intensivist / anaesthetist / physician / microbiologist / infectious diseases consultant (as required) with appropriate referral and retrieval is essential. Suspicion of severe sepsis or septic shock should trigger urgent referral to a tertiary centre with adult intensive care facilities.
> Follow the Surviving Sepsis Campaign Resuscitation ‘Bundle’ in severe sepsis.
> Monitor and maintain airway, breathing and circulation as first priority.
> In the event of hypotension and / or lactate > 4 mmol/L initiate urgent fluid resuscitation.
> Obtain blood cultures before antibiotic administration if possible but do not delay commencement of antibiotics.
> Systemically ill patients should commence broad spectrum intravenous antibiotics within 1 hour of recognition of severe sepsis.
> In a critically ill pregnant woman, birth of the baby may be considered if it would be beneficial to the mother or baby.
### Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ARDS</td>
<td>Acute respiratory distress syndrome</td>
</tr>
<tr>
<td>AVPU</td>
<td>Alert, verbal, pain, unresponsive</td>
</tr>
<tr>
<td>SaO₂</td>
<td>Arterial oxygen saturation</td>
</tr>
<tr>
<td>ASAP</td>
<td>As soon as possible</td>
</tr>
<tr>
<td>BP</td>
<td>Blood pressure</td>
</tr>
<tr>
<td>BSLs</td>
<td>Blood sugar levels</td>
</tr>
<tr>
<td>C</td>
<td>Celsius</td>
</tr>
<tr>
<td>CBP</td>
<td>Complete blood picture</td>
</tr>
<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>CVP</td>
<td>Central venous pressure</td>
</tr>
<tr>
<td>CVS</td>
<td>Cerebro-vascular system</td>
</tr>
<tr>
<td>EGDT</td>
<td>Early goal directed therapy</td>
</tr>
<tr>
<td>ECG</td>
<td>Electrocardiograph</td>
</tr>
<tr>
<td>GAS</td>
<td>Group A beta haemolytic streptococci</td>
</tr>
<tr>
<td>GCS</td>
<td>Glasgow coma scale</td>
</tr>
<tr>
<td>HCO₃⁻</td>
<td>Bicarbonate ion</td>
</tr>
<tr>
<td>HCT</td>
<td>Haematocrit</td>
</tr>
<tr>
<td>HVS</td>
<td>High vaginal swab</td>
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<tr>
<td>ICU</td>
<td>Intensive care unit</td>
</tr>
<tr>
<td>kg</td>
<td>Kilogram(s)</td>
</tr>
<tr>
<td>LFTs</td>
<td>Liver function tests</td>
</tr>
<tr>
<td>LVS</td>
<td>Low vaginal swab</td>
</tr>
<tr>
<td>MAP</td>
<td>Mean arterial blood pressure</td>
</tr>
<tr>
<td>mmHg</td>
<td>Millimetres of mercury</td>
</tr>
<tr>
<td>mL</td>
<td>Millilitre(s)</td>
</tr>
<tr>
<td>MSSU</td>
<td>Micro specimen of urine</td>
</tr>
<tr>
<td>PCO₂</td>
<td>Carbon dioxide partial pressure</td>
</tr>
<tr>
<td>PO₂</td>
<td>Oxygen partial pressure</td>
</tr>
<tr>
<td>PPROM</td>
<td>Preterm prelabour rupture of the membranes</td>
</tr>
<tr>
<td>RBC</td>
<td>Red blood cells</td>
</tr>
<tr>
<td>RCOG</td>
<td>Royal College of Obstetricians and Gynaecologists</td>
</tr>
<tr>
<td>ScvO₂</td>
<td>Central venous oxygen saturation</td>
</tr>
<tr>
<td>SpO₂</td>
<td>Oxygen saturation measured by pulse oximetry</td>
</tr>
<tr>
<td>SBP</td>
<td>Systolic blood pressure</td>
</tr>
<tr>
<td>SIRS</td>
<td>Systemic inflammatory response syndrome</td>
</tr>
<tr>
<td>WCC</td>
<td>White cell count</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organisation</td>
</tr>
</tbody>
</table>
Definitions

| Systemic Inflammatory Response Syndrome (SIRS) | More than one of the following clinical findings:
| o Temperature > 38°C or < 36°C
| o Heart rate > 90 per minute
| o Hyperventilation (evidence by respiratory rate > 20 per minute or PCO₂ < 32 mm Hg
| o WCC > 12,000 or < 3,000 |
| Sepsis | Presence of both infection (invasion of tissue, fluid or a body cavity by pathogenic micro-organisms) and systemic manifestations of inflammatory response syndrome (SIRS) |
| Severe Sepsis | Sepsis complicated by sepsis-induced organ dysfunction or developing tissue hypoperfusion |
| Septic shock | Persistence of hypoperfusion (hypotension) in a septic patient, despite adequate volume resuscitation
| o Hypotension is defined as: Systolic blood pressure (SBP) < 90 mm Hg or mean arterial blood pressure (MAP) < 60 mm Hg, or reduction of SBP > 40 mm Hg from baseline |
| Puerperal Sepsis | The term is used to describe sepsis occurring after delivery and the World Health Organisation (WHO) has defined it as “infection of the genital tract, occurring at any time between rupture of membranes or labour, and the 42nd day postpartum”, in which two or more of the following are present:
| > pelvic pain
| > fever
| > abnormal vaginal discharge
| > abnormal smell of discharge
| > delay in postpartum reduction of size of uterus |

Introduction

> Sepsis may arise in pregnancy at any time: before delivery, during labour or postpartum.
> Sepsis may arise from many sources and is not limited to infections arising from the genital tract
> The onset of sepsis may be insidious; women with severe sepsis may appear very well before suddenly collapsing, with little or no warning
> Early identification of severe sepsis allows prompt, appropriate multidisciplinary management
> Suspicion of severe sepsis or septic shock should trigger urgent referral to a tertiary centre with adult intensive care facilities
> Up to 20 to 30 % of intensive care unit (ICU) admissions of obstetric patients result from sepsis in pregnancy
> The contribution of sepsis as a cause of maternal mortality is between 3 % in developed countries and 12 % in developing countries
> One third of early maternal mortality is due to refractory hypotension. Late maternal mortality is due to multiple organ failure
> Urinary tract infection and chorioamnionitis are common infections associated with septic shock in the pregnant woman
> In the United Kingdom, the rapid course coupled with late presentation of some women with sepsis have contributed to a recent increase in direct maternal mortality due to genital tract sepsis (especially secondary to Group A streptococcal disease)
Common causes of sepsis in pregnancy

Common sites of infection in pregnancy
- Urinary tract (pyelonephritis)
- Pelvic structures (chorioamnionitis and endometritis)
- Surgical wounds (caesarean section, perineal laceration)
- Breast (mastitis)

Other causes
- Infection of intravenous cannula sites
- After urological procedures in the presence of urinary tract infection
- Related to regional anaesthesia e.g. spinal / epidural abscess (rare)
- Pneumonia (viral and bacterial)
- Intrauterine fetal death
- Septic abortion
- Acute appendicitis
- Acute cholecystitis
- Pancreatitis
- Bowel perforation (more common with inflammatory bowel disease)
- Necrotising fasciitis
- Meningitis

Common pathogens
- The most prevalent bacterial organisms responsible for severe infection include:
  - Group A beta haemolytic streptococci (GAS), also known as Streptococcus pyogenes
  - Group B streptococcus
  - Escherichia coli
  - Klebsiella
  - Staphylococcus aureus
  - Anaerobes: peptostreptococci, peptococci, bacteroides, clostridium
- Clostridium species and listeria monocytogenes are less common
- Viral causes e.g. influenza, varicella, hepatitis and herpes simplex
- Malaria and other tropical infections
Common bacterial infections and pathogens related to septic shock

<table>
<thead>
<tr>
<th>Infections and usual pathogens (in brackets – see right column)</th>
<th>Pathogens</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyelonephritis (1,4)</td>
<td>1. Escherichia coli</td>
</tr>
<tr>
<td>Pneumonia (6,7)</td>
<td>2. Bacteroides</td>
</tr>
<tr>
<td>Chorioamnionitis (1,2,8-12)</td>
<td>3. Clostridium</td>
</tr>
<tr>
<td>Endomyometritis (primarily after caesarean section) (1,2,5,9,12)</td>
<td>4. Klebsiella</td>
</tr>
<tr>
<td>Sepsis after miscarriage or termination of pregnancy (1,3)</td>
<td>5. Pseudomonas aeruginosa</td>
</tr>
<tr>
<td>Caesarean wound infection (1,2,6,7)</td>
<td>6. Streptococcus species</td>
</tr>
<tr>
<td>Severe mastitis (7)</td>
<td>7. Staphylococcus aureus</td>
</tr>
<tr>
<td>Necrotising fasciitis (2,3,6,9)</td>
<td>8. Group B streptococcus</td>
</tr>
<tr>
<td></td>
<td>9. Peptostreptococcus</td>
</tr>
<tr>
<td></td>
<td>10. Enterococcus</td>
</tr>
<tr>
<td></td>
<td>11. Listeria monocytogenes</td>
</tr>
<tr>
<td></td>
<td>12. Enterobacter</td>
</tr>
</tbody>
</table>

Risk factors for maternal sepsis

- Obesity
- Impaired glucose tolerance / diabetes
- Impaired immunity / immunosuppressant medication
- Anaemia
- Vaginal discharge
- History of pelvic infection
- History of group B streptococcal infection
- Amniocentesis and other invasive procedures
- Cervical Cerclage
- Prolonged spontaneous rupture of membranes
- Diabetes Mellitus
- Group A streptococcal (GAS) infection in close contacts / family members (working with or having young children)
- Of black or other minority ethnic group origin
- Retained products of conception
- Caesarean birth (greater risk associated with emergency caesarean)

Diagnosis of infection

A thorough history and physical examination is required to make the diagnosis of sepsis. Important elements in making a diagnosis include:

1. Having a suspicion of the diagnosis of sepsis
2. Assessing for evidence of end organ dysfunction
3. Identifying site / source of the infection
4. The speed of onset or deterioration in symptoms and signs is important

Symptoms

- Chills, sweating, warm skin, faintness or syncope, vomiting, rash, headache, dyspnoea, general weakness (or pain related to sites of sepsis – see below)
Signs

One or more of the following:

- Pyrexia (38°C or more): May not always be present and is not necessarily related to the severity of sepsis. Temperature may be less than 36°C in severe sepsis
- Tachypnoea (20 breaths or more per minute)
- Hypoxia
- Tachycardia (90 or more bpm)
- Arterial hypotension (systolic blood pressure less than 90 mm Hg; mean arterial pressure less than 70 mm Hg; or systolic blood pressure decrease 40 or more mm Hg)
- Decreased capillary refill or mottling
- Fetal distress secondary to maternal acidosis
- Oliguria (urine output less than 0.5 mL / kg / hr for at least two hours, despite adequate fluid resuscitation
- Considerable oedema or positive fluid balance (20 or more mL / kg over 24 hours)
- Ileus (absent bowel sounds)
- Impaired mental state, altered conscious level
- Hyperglycaemia in the absence of diabetes (plasma glucose more than 7.7 mmol / L)
- Bruising or discolouration of skin suggests late fasciitis (often pain receding as cutaneous anaesthesia supervenes as nerves die)
- Failure to respond to treatment

Assessment for end-organ dysfunction

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<thead>
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<th>End-organ dysfunction</th>
<th>Assessment</th>
</tr>
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<tbody>
<tr>
<td><strong>Acute circulatory failure</strong></td>
<td>Cardiovascular assessment: Peripheral perfusion, pulse, BP, CVP including signs of high or low cardiac output</td>
</tr>
<tr>
<td><strong>Metabolic acidosis</strong></td>
<td>pH, HCO₃, PaCO₂, lactate, anion gap</td>
</tr>
<tr>
<td><strong>Acute hypoxaemia</strong></td>
<td>SpO₂, PaO₂, PaO₂ / FIO₂</td>
</tr>
<tr>
<td><strong>Liver dysfunction</strong></td>
<td>LFTs</td>
</tr>
<tr>
<td><strong>Coagulopathy</strong></td>
<td>CBP and coagulation profile</td>
</tr>
<tr>
<td><strong>Acute renal failure</strong></td>
<td>Fluid overload, urine output, Urea, creatinine, electrolytes</td>
</tr>
<tr>
<td><strong>CNS disturbance</strong></td>
<td>Sedation score AVPU GCS Encephalopathy</td>
</tr>
<tr>
<td>e.g. altered mental state</td>
<td>Assessment for a hyper-catabolic state: negative nitrogen balance (e.g. low albumin) or raised blood glucose levels</td>
</tr>
</tbody>
</table>
Evaluate potential sources of infection

- Identify if any recent sore throat or respiratory illness or close contact with persons with illness (particularly streptococcal infections)
- Exclude intravenous drug misuse (high risk of staphylococcal and streptococcal sepsis)
- Recent febrile illness (especially if associated with chills and rigors - suggest bacteraemia or viraemia)
- Gastrointestinal symptoms (e.g. diarrhoea and vomiting) may be due to food borne pathogens or early toxic shock
- Consider need for vaginal examination to exclude retained tampons or swabs
- Exclude multi-resistant organisms such as ESBL-producing gram negative bacteria, vancomycin-resistant enterococci and MRSA (may affect choice of antimicrobial to combat sepsis and infection control precautions)
- Early investigation of non-specific symptoms in pregnant women is necessary to exclude serious infection.17

Investigations

<table>
<thead>
<tr>
<th>Blood tests</th>
<th></th>
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<tbody>
<tr>
<td>Serum lactate</td>
<td>Greater than 4 mmol/L requires urgent action</td>
</tr>
<tr>
<td>Complete blood picture</td>
<td>White blood cell (WBC) count &gt; 12 x 10^9 Leucopenia - WBC count &lt; 4 x 10^9 Normal WBC count with &gt; 10 % immature forms</td>
</tr>
<tr>
<td>Plasma C-reactive protein</td>
<td>More than 7 mg/L (usually significantly higher in bacterial sepsis)</td>
</tr>
<tr>
<td>Urea and electrolytes</td>
<td>Creatinine rise of &gt; 44.2 micromol/L; sepsis is severe if creatinine level &gt; 176 micromol/L</td>
</tr>
<tr>
<td>Plasma glucose</td>
<td>Hyperglycaemia in the absence of diabetes (plasma glucose &gt; 7.7 mmol/L)</td>
</tr>
<tr>
<td>Liver function tests (LFTs)</td>
<td>Hyperbilirubinaemia (plasma total bilirubin &gt; 70 micromol/L)</td>
</tr>
<tr>
<td>Coagulation profile</td>
<td>Coagulation abnormalities (INR &gt; 1.5 or APTT &gt; 60 seconds)</td>
</tr>
<tr>
<td>Blood gas</td>
<td>Arterial hypoxaemia (PaO_2 / FIO_2 &lt; 300 mm Hg) Sepsis is severe if &lt; 250 mm Hg in the absence of pneumonia or &lt; 200 mm Hg in the presence of pneumonia Raised serum lactate ≥ 4 mmol/L</td>
</tr>
</tbody>
</table>

Adapted from: RCOG Diagnostic criteria for sepsis (modified from Levy⁶)
Microbiology

> Blood cultures (ideally obtain before antibiotic administration)
> Obtain cultures of additional sites as indicated and as soon as possible e.g. midstream specimen urine (MSSU), wound swab (episiotomy or caesarean section), placental swabs, respiratory secretions, naso-pharyngeal aspirate (NPA), amniotic fluid, cerebrospinal fluid, HVS, LVS, endocervical, expressed breast milk
> If the methicillin-resistant Staphylococcus aureus (MRSA) status is unknown, obtain swabs from nose, groin and axilla and send for urgent screening
> Suspected pneumonia - Chest X-ray, naso-pharyngeal aspirate

Imaging

> Consider imaging modalities e.g. Chest X-ray, pelvic ultrasound, computed tomography, magnetic resonance imaging (may help define inflammation or the collection of pus)
> Echocardiography may be useful in cases of women who are intravenous drug users, particularly those with staphylococcal bacteraemia as there may be right-sided endocarditis and to assess cardiac function

Site of infection – specific considerations

Early diagnosis to determine if surgical intervention required
May be difficult to localise the source

Chorioamnionitis

Diagnosis of chorioamnionitis relies on the clinical presentation
> Maternal fever > 38° C with any 2 of the following:
> Increased white cell count (>15 x 10⁹/L)
> Maternal tachycardia (> 100 bpm)
> Fetal tachycardia (>160 bpm)
> Uterine tenderness
> Offensive smelling vaginal discharge
> C-Reactive Protein > 40
> Histological examination of placenta and membranes with evidence of acute inflammation may confirm diagnosis after birth

Treatment of chorioamnionitis
> Amoxicillin 2 g IV every 6 hours
> Gentamicin 5 mg/kg IV daily
> Metronidazole 500 mg IV every 12 hours
> If allergic to penicillin, give clindamycin 450 mg IV in 50 – 100 mL over at least 20 minutes every 8 hours AND gentamicin 5 mg /kg IV daily until delivery
> Do not inhibit labour, but consider hastening delivery under intravenous antibiotic cover
> Consider optimal mode of delivery (LSCS versus vaginal birth) on the basis of the findings and the anticipated duration until birth
> Consider ongoing treatment with antibiotics postnatally

Endometritis

> Offensive lochia
> Unusual bleeding
> Consider need for evacuation of retained products of conception
> If does not respond to antibiotics, consider septic pelvic thrombosis and commence on heparin
Urinary tract infection
> Loin pain
> Tenderness to renal percussion
> Send midstream specimen of urine (MSSU)
> Treat acute pyelonephritis aggressively (For more information see ‘Urinary tract infections in pregnancy’ in the A to Z index at www.sahealth.sa.gov.au/perinatal)

Mastitis
> Reddened sector in the breast
> Immediate referral to hospital if the woman is clinically unwell, no response to oral antibiotics within 48 hours, mastitis recurs or there are severe or unusual symptoms
> Complications include breast abscess, necrotising fasciitis, toxic shock syndrome

Wound, skin and soft tissue infections
> Signs of inflammation at the site
> Fever, chills, tachycardia, tachypnoea, wound swelling, warm to touch, painful, discharge
> Closed space infections need surgical drainage
> Carefully examine intravenous cannulae or injection sites, and caesarean or episiotomy wounds
> Take a swab of any discharge
> Exclude complications of venous thrombosis or necrotising fasciitis (see below)

Necrotising fasciitis
> Extreme pain, associated with signs of prostration and severe sepsis
> Requires early surgical intervention with fasciotomy and aggressive antibiotic treatment

Bacteraemia and septic shock
> Suspected bacterial sepsis requires careful examination for skin and soft tissue infection
> If drains, vascular access devices or other indwelling devices are suspected as the source of infection, remove as soon as is practicable
> Skin and soft tissue infections are particularly associated with toxic shock syndromes
> Septicaemic seeding of streptococci from a uterine focus may give rise to a secondary focus in a limb, simulating a venous thrombosis
> Women with thrombosis who are systemically unwell with any features of sepsis should be examined very carefully. If shock or other organ dysfunction present, rapid referral to adult intensive care

Pneumonia
> Fever, chills, dyspnoea, cough, sputum production, pleuritic pain
> Consider referral to respiratory physician, medical microbiologist
> Send sputum sample, throat swab for culture
> Chest X-ray

Gastroenteritis
> Diarrhoea and vomiting may be symptoms of bacterial sepsis, due to organisms including staphylococcus and streptococcus
> Stool sample for routine culture (include C. difficile toxin testing if diarrhoea offensive after antimicrobial treatment)
> Salmonella and Campylobacter rarely cause severe systemic infection
Pharyngitis
- Usually viral
- Approximately 10% of cases are due to GAS
- Obtain throat swab for culture
- Antibiotic treatment if 3 or more of the following: fever, tonsillar exudate, no cough, tender anterior cervical lymphadenopathy

Infection post termination of pregnancy / miscarriage
- Suspect infection in women with a history of a recent termination of pregnancy or spontaneous miscarriage who present with pyrexia, persistent bleeding or abdominal pain, especially if the pain is constant and severe
- Vaginal swabs, ultrasound scan to exclude retained products of conception and consider diagnostic evacuation of uterus (evacuation of retained products of conception) if there is still doubt
- Complete blood picture, C-reactive protein, blood cultures if pyrexia > 38°C Celsius
- Commence high dose broad-spectrum intravenous antibiotics without waiting for microbiology results

Cholecystitis / Cholangitis
- Upper right quadrant pain, biliary colic after a fatty meal, jaundice, nausea, vomiting, fever, ultrasonography examination (USG) of abdomen

Epidural abscess
- Very rare complication
- S. aureus is the usual organism
- Severe backache
- Neurological deficit including bowel / bladder dysfunction and unexplained fever

Management
- After identification of severe sepsis, in the first 6 hours, follow the Surviving Sepsis Campaign Resuscitation ‘Bundle’ (see below)
- Treat underlying infection and support failing organ functions
- Systemically ill patients should commence broad spectrum, intravenous antibiotics (within 1 hour of recognition of severe sepsis)
- Patients with single organ failure not responding to simple measures require transfer to a tertiary centre with high dependency care
- Patients with two or more organ failures and respiratory failure require transfer to a tertiary centre with Adult Intensive Care Facilities (see ‘Assess for evidence of end-organ dysfunction’ above)
- Key Practice Point for outpatients:
  Abdominal pain, fever (> 38°C), tachypnoea and sustained tachycardia (> 90 beats per minute) are indications for admission and intravenous antibiotics
Surviving Sepsis Campaign Resuscitation ‘Bundle’

<table>
<thead>
<tr>
<th>Tasks to be performed within the first 6 hours of the identification of severe sepsis7</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt; Obtain blood cultures before antibiotic administration</td>
</tr>
<tr>
<td>&gt; Administer broad spectrum antibiotic within one hour of recognition of severe sepsis</td>
</tr>
<tr>
<td>&gt; Measure serum lactate</td>
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<tr>
<td>&gt; In the event of hypotension and / or lactate &gt; 4 mmol/L:</td>
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<tr>
<td>&gt; Deliver an initial minimum 20 mL / kg of crystalloid or an equivalent fluid</td>
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<tr>
<td>&gt; Apply vasopressors for hypotension not responding to initial fluid resuscitation to maintain mean arterial pressure above 65 mm Hg</td>
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<tr>
<td>&gt; In the event of persistent hypotension despite fluid resuscitation (septic shock) and / or serum lactate &gt; 4 mmol/L:</td>
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<tr>
<td>&gt; Achieve a central venous pressure (CVP) of ≥ 8 mm Hg</td>
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<tr>
<td>&gt; Achieve a central venous oxygen saturation (ScvO2) ≥ 70 % or mixed venous oxygen saturation (ScvO2) ≥ 65 %</td>
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</tbody>
</table>

Adapted from the Surviving Sepsis Campaign Resuscitation Bundle (group of therapies)15

Initial management (0-6 hours)

> Early involvement of senior obstetrician / intensivist / anaesthetist / physician / microbiologist / infectious diseases consultant (as required) with appropriate consultation, referral and retrieval

> Assess need for central venous access and initiate Surviving Sepsis Campaign Resuscitation Bundle if severe sepsis is suspected (see table 3 above)

> Follow the A, B, C, D, E’s of resuscitation

Airway

> Maintain a clear airway

> Administer oxygen via non-rebreathing mask at 12 litres per minute

> If the woman has collapsed, check her airway is patent, she is breathing and maintain her position in left-lateral

Breathing

> Assess breathing pattern, rate and colour and ventilate if required

> Attach pulse oximeter, electrocardiogram and automatic blood pressure monitors and monitor SpO2, maternal pulse and blood pressure

Circulation

> Hypovolaemia is present in almost all patients with septic shock

> Obtain intravenous access

> Take blood for blood cultures, serum lactate, complete blood count, coagulation profile, urea and electrolytes, liver function tests, blood glucose

> Immediate fluid resuscitation with either crystalloid or colloid as per medical order

> If no response to simple measures of fluid resuscitation consider the need for insertion of central venous catheter and CVP monitoring

> Correct hypotension – Administer vasopressors for hypotension that is not responding to initial fluid resuscitation to maintain mean arterial pressure (MAP) more than 65 mm Hg (intensivist or anaesthetic management)

> Perform arterial blood gases
Documentation and differential diagnosis

> Obtain history as soon as possible
> Document vital signs on ‘Rapid deterioration and Response’ chart
> Confirm diagnosis
> Commence broad spectrum intravenous antibiotic treatment within 1 hour of diagnosis of severe sepsis
> Commence fluid balance chart and monitor urine output

NB: Fluid overload may lead to fatal pulmonary or cerebral oedema in women with septic shock. Clear, accurate documentation and careful monitoring of fluid balance is essential

Fetal wellbeing

> Assess and monitor fetal wellbeing (cardiotocography) if applicable
> Continuous fetal monitoring intrapartum
> Consider the need for delivery as applicable
  > In a critically ill pregnant woman, birth of the baby may be considered if it would be beneficial to the mother or the baby or to both. A decision on the timing and mode of birth should be made by a senior obstetrician after discussion with the woman if her condition allows
  > If preterm birth is anticipated, cautious consideration should be given to the use of antenatal corticosteroids for fetal lung maturity in the woman with sepsis
  > Epidural / spinal anaesthesia should be avoided in women with sepsis

Remove the source of the maternal infection

> Expedite delivery if there are signs of chorioamnionitis
> Any retained products of conception should be removed as soon as the maternal condition is stable
> If large amounts of pelvic pus are present a laparotomy and sometimes hysterectomy may be necessary

Maternal observations

> Obtain and document temperature, pulse, respiratory rate, blood pressure, SpO₂ every 15 minutes in the initial assessment phase
  > Once stable, continue hourly
> Utilise ‘Rapid Deterioration and Response’ chart. The use of early warning charts and escalation guidelines including involvement of senior medical staff (intensivist, physician) assists in the early detection and management of the deteriorating patient
> Glasgow Coma Scale and pupil response if required
> Hourly urine output (consider insertion of indwelling catheter)
> In rapidly deteriorating cases, ensure urgent referral to critical care team and obstetric consultant
Antibiotics

> Early consult with microbiologist or infectious diseases consultant for women with evidence of systemic infection
> Commence broad spectrum intravenous antibiotic cover within 1 hour of suspicion of severe sepsis, with or without septic shock
> The choice of antibiotic depends on the clinical suspicion, local flora and culture information, if available
> If genital tract sepsis is suspected, prompt early treatment with a combination of high-dose broad-spectrum intravenous antibiotics may be lifesaving
> Empirical treatment should include broad spectrum antimicrobials active against Gram-negative bacteria, and capable of preventing exotoxin production from Gram-positive bacteria (according to local microbiology policy). Gram-positive cover is necessary if the likelihood of this infection is high

Thromboprophylaxis

> Women with sepsis are at increased risk of venous thromboembolism
> Consider prophylaxis with low molecular weight heparin
> Use of compression stockings

### Staphylococcal and streptococcal toxic shock syndrome

Toxic shock syndrome is a life-threatening illness characterised by hypotension and multi-organ failure. It may be caused by Staphylococcus aureus (rarely isolated) or Streptococcus pyogenes that produce and release superantigenic exotoxins. The exotoxins activate T-cells on a large scale resulting in a massive release of inflammatory cytokines.

<table>
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<tr>
<th>Staphylococcal toxic shock syndrome</th>
<th>Streptococcal toxic shock syndrome</th>
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| 1. Fever ≥ 39.9°C | A. Isolation of group A Streptococcus from:
2. Rash: diffuse macular erythema | 1. Normally sterile site: blood, cerebrospinal fluid, peritoneal fluid, tissue biopsy
3. Desquamation: 10-14 days after onset of illness, especially palms and soles | 2. Non-sterile site: throat, vagina, sputum
4. Hypotension: systolic BP < 90 mmHg |
5. Multisystem involvement: 3 or more of the following systems affected: |
  o GIT: vomiting or diarrhoea at onset of illness |
  o Muscular: severe myalgia or elevated creatinine phosphokinase |
  o Mucous membranes: vaginal, oropharyngeal or conjunctival hyperaemia |
  o Renal: creatinine twice the upper limit of normal |
  o Haematological – platelets ≤ 100 x 10⁹/L |
  o Central nervous system – disorientation or alterations in consciousness without focal neurological signs |
| Case classification: |
  o **Probable**: 4 of the 5 clinical findings positive |
  o **Confirmed**: case with all 5 clinical findings |
  o **Probable**: meets clinical case definition (above) plus isolation from non-sterile site |
  o **Definite**: meets clinical case definition (above) plus isolation of group A Streptococcus from a normally sterile site |
Intravenous immunoglobulin (IVIG)

- IVIG is recommended for severe invasive streptococcal or staphylococcal infection if other therapies have failed
- High dose IVIG has been used in pregnant and postpartum women and is effective in exotoxic shock (i.e. toxic shock attributable to streptococci and staphylococci)\(^8\)
- For more information, see ‘Intragam infusion’ in the A to Z index at [www.sahealth.sa.gov.au/perinatal](http://www.sahealth.sa.gov.au/perinatal)

Complications

- Acute respiratory distress syndrome (ARDS)
- Disseminated Intravascular coagulation
- Renal failure
- Hepatic failure
References


Useful online references


Surviving Sepsis Campaign at http://www.survivingsepsis.org
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