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FOREWORD

Sepsis is a world-wide public health issue and claims thousands of lives each year. The incidence of sepsis is escalating as the population ages, and its treatment is becoming an increasingly significant burden on national health care expenditure.

Sepsis is a medical emergency just like a heart attack or stroke. It arises when the body’s response to infection causes a generalised inflammatory response that can lead to shock, multiple organ failure and death. Early recognition and prompt treatment is essential to improve survival.

The Clinical Excellence Commission (CEC) launched the SEPSIS KILLS program in May 2011 in emergency departments and the program has been extended to the inpatient wards. The program seeks to engage and support our clinical staff in hospitals across New South Wales to make sustainable improvement in the recognition and management of severe infection and sepsis.

Patients with sepsis are high risk and may deteriorate rapidly. The program is closely aligned with the Between the Flags system which facilitates early recognition, response and management of the deteriorating patient.

Sepsis awareness, early identification, resuscitation and referral to specialist care will ensure that all patients with sepsis receive timely and appropriate care. I encourage each of your facilities, and all of your staff (clinical and managerial) to implement the program.

Clifford F Hughes, AO
CLINICAL PROFESSOR
CEO, CLINICAL EXCELLENCE COMMISSION
INTRODUCTION

Welcome to the Inpatient SEPSIS KILLS program implementation toolkit. It is a practical guide for managers and clinicians to improve the recognition and management of sepsis in adult and paediatric patients in inpatient wards of medium and large hospitals in NSW.

It has been developed from the collective knowledge and experience of clinicians and CEC staff who have been involved in the SEPSIS KILLS program development and implementation.

The toolkit provides:
- Tools to plan, implement and evaluate a sepsis improvement project in hospital wards
- Education resources
- Communication resources

The toolkit is not exhaustive. Over time, new tools and resources will be developed in response to clinician feedback and program needs. These resources will be made available on the SEPSIS KILLS program page of the CEC website www.cec.health.nsw.gov.au/programs/sepsis.
BACKGROUND

Sepsis is one of the leading causes of death in hospital patients worldwide. It causes more deaths than prostate cancer, breast cancer and HIV/AIDS combined\(^1\). There are approximately 15,000 cases of severe sepsis and septic shock in Australian and New Zealand intensive care units per year\(^2\).

Sepsis is a life-threatening condition that arises when the body’s response to infection injures its own tissues and organs\(^3\). Sepsis can present in any patient, in any clinical setting and is a medical emergency. Despite this, awareness of sepsis and the need for prompt and targeted treatment is limited.

Appropriate recognition and timely management of patients with severe infection and sepsis is a significant problem in NSW hospitals and in health care facilities around the world. Delayed treatment is associated with high mortality rates, significant morbidity and high costs to the health care system. In paediatric patients, sepsis is one of the leading causes of death, with mortality rates as high as 10 per cent. Many of these deaths are preventable\(^4\).

The mortality rate for adult patients with septic shock is around 25 per cent\(^5\). It has been shown to increase by 7.6 per cent for every hour of delay, after the onset of hypotension, in starting antibiotic therapy\(^6\). In the complex hospital ward environment, there are frequently long delays between medical review and antibiotic prescription, particularly when decision making is by junior medical officers\(^7\).

The CEC Clinical Focus Report on Recognition and Management of Sepsis\(^8\) found deficits in a range of clinical settings in NSW health care facilities in 2009. The lack of timely recognition and appropriate management was further demonstrated in the findings from the 2011 CEC Quality Systems Assessment\(^9\).

The CEC SEPSIS KILLS Economic Analysis\(^10\) estimates that if the status quo is maintained over the next ten years, sepsis-related conditions in the NSW health system will constitute $3.7 billion, 1.3 million bed days, 701,000 cost weighted separations and an unknown number of potentially avoidable deaths.
SEPSIS KILLS PROGRAM

The SEPSIS KILLS program is a quality improvement initiative that has been developed from international evidence-based practice\textsuperscript{11,12,13}. The goal is to reduce preventable harm to patients with sepsis through early recognition and prompt management.

It provides significant benefits at both the clinical and system levels, including:

- Enhanced clinician skills in sepsis recognition and management
- More timely, standardised and effective detection and management of sepsis
- Reduced mortality, morbidity and bed-stays from sepsis-related conditions
- Improved quality and safety of care and a better and safer patient experience.

The program started in May 2011 and has been implemented in 180 NSW public hospital emergency departments. Since it was launched, time to antibiotic administration has greatly improved. Median time to antibiotic administration in emergency departments has been reduced from more than four hours, to consistently less than sixty minutes in 2013 and 2014.
SEPSIS AND THE DETERIORATING PATIENT

The inpatient phase of the SEPSIS KILLS program is focused on improving the recognition and management of sepsis in adults and children in the inpatient wards of public hospitals in NSW. It directly links with the Between the Flags system (BTF), the NSW Health Policy: Recognition and Management of Patients who are Clinically Deteriorating (PD2013_049) and the Australian Commission on Safety and Quality in Health Care national accreditation standards\textsuperscript{14}.

Deteriorating patients with sepsis can be escalated via the NSW Standard Observation Charts and the Clinical Emergency Response System (CERS). By undertaking routine clinical observations, nurses and doctors play a vital role in identifying and escalating sepsis. The clinician is encouraged to think why the patient is deteriorating and ask “could this be sepsis?”.

The ‘track and trigger’ design of the NSW Health Standard Observation Charts includes standard calling criteria for Clinical Review and Rapid Response which are pivotal in supporting clinical staff to recognise deterioration.

All NSW public hospitals and acute care facilities have an established CERS so that deteriorating patients receive an appropriate clinical response. CERS are developed and tailored to the health service/facility’s needs and resources.

The main components of this system of escalation include:

- A Clinical Review process that has the capacity to respond within 30 minutes to a breach in Yellow Zone observations or additional criteria
- A Rapid Response process that is immediately available in response to a breach in the Red Zone observations or additional criteria
- Necessary equipment is available to perform advanced resuscitation
- An escalation process for transferring patients that require a higher-order of care to a facility that can provide it.

Preliminary NSW sepsis data suggests that 30 per cent of adults who require a Rapid Response are septic. This is replicated in national and international literature\textsuperscript{15}, with sepsis being a leading cause for clinical deterioration, accounting for one in three calls for a Rapid Response team.

The tools provided by the BTF system and SEPSIS KILLS program give clinicians objective criteria, backed by policy for escalation, which empowers them to call for senior clinical help when they judge a patient to be deteriorating or are concerned about the patient’s condition. This is illustrated in the following diagram:

1. Recognition
   - Is my patient ‘Between the Flags’? (BTF)

2. Response
   - If not, what should I do? (Yellow & Red Zone response)

3. Root Cause
   - Why is my patient deteriorating? (SEPSIS KILLS)

4. Response
   - How should I treat him or her? (SEPSIS KILLS)
FIVE ELEMENTS OF THE PROGRAM

The five elements of the SEPSIS KILLS program align with the BTF system and are designed to establish a sustainable Statewide initiative which will deliver improved systems for recognition and response to patients with sepsis.

The five inter-dependent elements of the program are:

1. Governance: establishment of guidelines for an administrative structure to oversee the implementation and sustainability of the system in the acute hospitals in NSW.


3. Sepsis Tools: to guide clinical decision making in parallel with the local CERS to facilitate the following:
   - RECOGNISE risk factors, signs and symptoms of sepsis
   - RESUSCITATE with rapid intravenous fluids and antibiotics within the first hour of recognition of sepsis
   - REFER to senior clinicians and specialty teams, including retrieval as required

4. Education: to ensure appropriate skills and knowledge for the recognition and management of the patient with sepsis.

5. Evaluation: standard performance indicators to be collected and used to inform users of the system and to guide sepsis recognition and management improvement.

The following sections of the Sepsis Toolkit provide guidelines, tools and resources to support implementation of the five elements in your health care facility.
GOVERNANCE

The success and long-term sustainability of the SEPSIS KILLS program is dependent on appropriate governance structures and processes at all levels of the health care facility. Leadership by both management and clinicians will enable a top-down, bottom-up approach to drive and sustain improved outcomes for patients with sepsis.

The governance of the program should be integrated with existing structures for the recognition and management of patients who are clinically deteriorating.

Key roles in the local health district (LHD)
Each LHD and health care facility will need to identify and appoint key position holders with operational responsibility for implementation of the program. These position holders will work closely with appropriate advisory committees, program teams and expert advisors. Suggested roles and responsibilities for LHD and facility personnel are detailed below.

**LHD executive sponsor**
- Ensure key personnel have been identified and appointed
- Establish an effective LHD governance structure which integrates with the deteriorating patient strategy
- Implement the five elements of the program in all medium and large LHD facilities
- Assist and support the LHD sepsis leads to coordinate the LHD implementation strategy.

**LHD sepsis lead**
- Coordinate the implementation of the program in all medium and large LHD facilities in collaboration with the LHD and facility executive sponsors
- Ensure stakeholders at LHD and facility levels have been consulted in the development of the implementation strategy
- Work with the facility executive sponsors to engage senior clinicians and department heads
- Liaise with the CEC on LHD progress and strategies developed to overcome barriers

**Facility executive sponsor**
- Appoint a facility sepsis program lead
- Establish an implementation team
- Establish an effective governance structure for the facility which integrates with the deteriorating patient strategy
- Identify and allocate resources to support implementation
- Ensure that all key personnel have been identified and appointed including medical and nursing ward/unit clinical leads/champions
- Assist and support ward/unit clinical leads/champions by endorsing the program as a vital initiative which is part of the deteriorating patient strategy
- Facilitate senior medical staff discussion on supervision and escalation processes for junior medical staff
- Provide reports to the LHD sepsis lead and executive sponsor on implementation progress and results

**Facility sepsis program lead**
- Assist and support ward/unit clinical leads/champions to engage senior clinicians and implement the sepsis program in their local areas
- Facilitate senior medical staff discussion on supervision and escalation processes for junior medical staff
- Coordinate facility sepsis communication and education strategies
• Ensure materials are available and promoted
• Provide regular feedback and progress reports to the facility Executive Sponsor

Facility clinical leads/champions
• Participate in the development of local systems for the program
• Work with the facility sepsis program lead and executive sponsor to engage senior clinicians and implement the program in wards/units
• Coordinate ward/unit based communication and education
• Support clinicians as they adapt to the altered clinical environment
• Provide feedback and progress reports to the facility sepsis program lead
Getting started
Each health care facility will need to develop a strategy to effectively implement the program in all wards. This section of the toolkit provides information on a ten-step implementation process which can be adapted to suit local environments.

STEP 1: Establish a case for change
Making a change to existing practice can be challenging and requires time, effort and resources. A good case for change will help to convince others that it is necessary to change the current practices i.e. it provides compelling evidence of why there is a problem, what will improve and the expected outcome.

A short written document and/or presentation can be used to gain people’s interest. Include evidence that timely sepsis recognition and management is an international problem and sepsis is a medical emergency. Local data showing how sepsis is currently managed is a powerful tool to convince clinicians why the change is necessary.

Gaining leverage from the outcomes of the emergency department sepsis project will also be influential. Benefits of the change need to be clearly indicated along with linkages to other quality and safety initiatives such as national accreditation requirements. The benefits may vary according to the target group e.g. clinical or managerial.

Resources
- Program summary
- Memo template
- Sepsis program general presentation

STEP 2: Establish governance arrangements and a program team
Governance is a critical foundational element of the program. Defining clear roles and responsibilities for governance is a high priority and need to be established early.

The program should be linked with the facility quality and safety plan, deteriorating patient strategy and relevant national accreditation standards.

The following roles need to be identified at both LHD and facility levels:
- Executive sponsor
- Program lead
- Program team to oversee the improvement process

The program team will need multidisciplinary senior managers and clinicians to provide organisational and ward/unit leadership and support. A responsible committee will need to be identified with agreement on communication and reporting channels between the program team, responsible committee and the LHD sepsis lead/clinical governance unit.

An initial task for the team is to develop an overall implementation plan. The CEC has developed an implementation plan template for NSW facilities to promote consistency and to help identify key elements required.

The plan will guide the team and ensure relevant activities are managed from the outset including risk assessment, communication and evaluation strategies. The program plan will be endorsed by the facility executive sponsor and the LHD sepsis lead.

Resources
- Facility implementation plan
- Facility implementation checklist
STEP 3: Recruit respected and influential leaders and champions
Senior leadership is crucial to the success of the program. It is recommended that each ward/unit has medical and nursing leads/champions. These people need to be senior clinicians who are willing and able to drive organisational change.

The sepsis leads from the emergency department will be able to provide helpful insights to the leadership role as well as ideas for the program implementation strategy. Some of the medical and nursing leads may also be program team members.

STEP 4: Identify sepsis management barriers and enablers
A review of the current process for managing sepsis should be undertaken. If resources are available, it is useful to undertake a retrospective audit of sepsis cases to provide insights into current practice.

A brainstorming session to identify the causes of inadequate or delayed recognition and treatment of sepsis will be helpful to better understand sepsis management barriers and enablers. Areas to consider are people, patients, processes, knowledge, skills, equipment, communication, escalation and supervision.

A list of common causes of delay in recognition and treatment of sepsis is provided in Section B: Tools and Resources. The list was derived from CEC pilot study feedback in several large hospitals. It should be noted however, that the brainstorming activity is far more effective if it is undertaken by the facility program team and by wards/units to identify the local issues.

When undertaking the brainstorming process, it is important to involve representatives from all levels/disciplines to ensure that relevant issues are identified. The program team or ward/unit can then develop an action plan to address the main barriers.

Resources
- Common causes of delay in recognising and treating sepsis in the wards
- Implementation action plan

STEP 5: Adapt the sepsis tools to align with the clinical emergency response system (CERS) and other local processes
A range of tools have been developed to support sepsis recognition and management. The CERS and Sepsis Tools section in the Toolkit provides details on each of the tools.

It is important to ensure that the sepsis pathway is aligned with the local CERS. This may include changing terminology if hybrid systems are in place. Discussions with senior medical staff should be facilitated to establish processes for escalation of sepsis to the Attending Medical Officer (AMO), infectious diseases experts and intensive care. Suggested triggers for escalation to the AMO and infectious diseases are provided in the Toolkit Section B to assist each department or facility to initiate discussions and establish local processes.

The CEC sepsis antibiotic guideline can be reviewed by the facility and/or LHD drug committee and aligned with local drug and antimicrobial stewardship processes.

Patients with sepsis are at a high risk of deterioration in the days following initial treatment. It is important that a high level of vigilance and monitoring is maintained. Local minimum monitoring requirements for ward patients with sepsis should be established and the CEC 48 hour sepsis management plan can be adapted for local use.
STEP 6: Identify tools and resources to support sepsis management
Other tools and resources may be required to support implementation of improved sepsis management. Sepsis trolleys or packs that hold the necessary equipment for managing a patient with sepsis have been found to be useful in both the emergency and ward settings. Sepsis trolleys also provide a clear message to ward staff that sepsis is an emergency and necessitates rapid intervention.

Establishing processes for timely lactate testing is important to support identification and management of sepsis. High serum lactate levels are strongly associated with increased mortality. Accessing results for formal serum lactate levels may take considerable time in most hospitals. Point of care testing may be available through the intensive care unit or emergency department and could be considered as an alternative process to provide quick access to lactate results.

Resources
- Ward implementation checklist
- Sepsis trolley/pack content list

STEP 7: Develop an education strategy
Training and education is important to initiate a clinical practice change and is also vital to help sustain and spread the change over time. An initial step is to identify the education tiers or groups and to document a plan to provide tailored education for specific professional and craft groups.

The strategy will vary in each facility however it may be helpful to consider how the education was rolled out for the BTF system. At a minimum, there will need to be awareness training (key sepsis messages and orientation to the sepsis pathway) for all nursing and medical staff in direct contact with patients. More detailed training can be provided for the Rapid Responders and other key stakeholders who will be required to provide clinical decision support to wards and clinicians.

Master classes for medical staff, nurse unit managers, educators, clinical nurse consultants and nurse managers could be adopted. Evaluation of the education strategy can be undertaken using the multiple choice questionnaire provided in Part 2: Tools and Resources or by using a locally developed pre and post education survey.

Education can be incorporated into the BTF Tier Two education (DETECT training), as well as orientation programs for junior medical officers and new staff starting at the hospital.

Resources
- SEPSIS KILLS video
- SEPSIS KILLS general presentation
- SEPSIS KILLS clinical presentation: adults
- SEPSIS KILLS clinical presentation: paediatrics
- SEPSIS KILLS e-learning module
- SEPSIS KILLS multiple choice questions
STEP 8: Pilot the new sepsis process
The program team will now be ready to implement a pilot study of the new sepsis process using short Plan, Do, Study, Act cycles. It is important to keep the pilot study a manageable size choosing one or two wards where there is strong clinical leadership and commitment to change. Groundwork will need to be done to prepare the staff in the pilot area with education and promotion activities.

Collecting feedback from stakeholders frequently is important. It is recommended that an issues log is used to document concerns as they arise (particularly important out of hours) as well as having regular forums to discuss progress. Reviewing and revising the process during and after the pilot will help to identify unintended consequences and reinforce the importance of staff feedback and generation of solutions.

Data collection to demonstrate improvement will need to be established at the outset of the pilot and should include quantitative data (time to IV fluid and antibiotic administration) as well as qualitative data (staff feedback).

Resources
Sepsis issues log

STEP 9: Sustain and spread the process to other wards/units
When the pilot study is complete, the process can be spread to other wards and units in the hospital in a phased approach. The pilot outcomes can be communicated using a variety of methods including meetings, newsletters and intranet posts to share the benefits gained and the lessons learnt.

Organizational guidelines on sepsis management principles and procedures can be developed and embedded into structures and routines e.g. clinical handover, orientation programs, junior medical staff education programs and BTF Tier Two education (DETECT training).

STEP 10: Measure, evaluate and improve
Evaluation is a systematic process to determine if an improvement is being made. Developing an evaluation plan is important for both the short term and long term success of an improvement program. Evaluation can help to identify any unintended consequences as well as providing lessons for future improvement.

It is suggested that consideration is given to using a range of evaluation methods. These could include medical record audit, observational audit, survey and focus groups. Displaying evaluation data and discussing the findings at ward meetings will help to drive improvement.

Each health care facility and LHD will establish processes for reporting results to local committees and the LHD. Further details on sepsis measures and methodology can be found in the Evaluation section of the Toolkit.
CERS AND SEPSIS TOOLS

A range of tools and resources are available to assist clinicians to identify and treat sepsis, in accordance with national and international best practice. The tools have been developed in collaboration with clinical experts and other key stakeholders from tertiary, metropolitan and regional/rural facilities across NSW.

The main three clinical tools are:
- Sepsis pathways
- Antibiotic guidelines
- Sepsis 48 hour management plan

The tools and resources are provided in Part B of the Toolkit and/or the sepsis page of the CEC website.

Sepsis pathways

The sepsis pathways for adult and paediatric inpatients provide clear guidelines for sepsis recognition, notification, escalation and initial management. They are built around three key actions:

- **RECOGNISE** risk factors, signs and symptoms of sepsis
- **RESUSCITATE** with rapid intravenous fluids and antibiotics within the first hour of recognition of sepsis
- **REFER** to senior clinicians and specialty teams including retrieval as required

The pathways are based on the following logic which directly links the pathways with the BTF system and the NSW Health Policy: Recognition and Management of Patients who are Clinically Deteriorating (PD2013_049).

The left side of the flow diagram illustrates the process used from recognition to treatment of patients with sepsis. The right side of the flow diagram describes the tools, processes and steps currently used in the hospital inpatient wards.

The following section provides information on the linkage of the logic (above) with the sepsis pathways.
Recognition
Recognition of sepsis risk factors, signs and symptoms may be facilitated via any of the following systems and relates to the recognition section of the sepsis pathway:

NSW Health Standard Observation Charts
In line with NSW Health policy, observations must be recorded on a NSW Health Standard Observation Chart (SAGO, or age-specific SPOC). In the absence of a documented monitoring plan, frequency of observations should be completed as indicated by the patient’s condition. This should be no less than three times per day, at eight hourly intervals.

REACH (Recognise, Engage, Act, Call, Help is on its way)
This is a patient and family activated Rapid Response program. REACH empowers patients and families to escalate care if they are concerned about the condition of the patient by first encouraging engagement with the treating clinicians at the bedside.

Ward Rounds
A patient may be identified as being at risk of or having sepsis during a ward round where medical, nursing and allied health staff comes together at the patient’s bedside to discuss the patient’s current clinical status and plan of care.

Escalation
A Clinical Review or Rapid Response must be activated for the deteriorating patient in accordance with local CERS. Patients who are identified as septic at the time of a ward round or during a routine medical consultation require a Rapid Response call if the patient’s observations are in the Red Zone.

Could this be sepsis?
Consideration should be given to whether sepsis or any other time critical conditions such as a new arrhythmia, hypovolaemia/haemorrhage, PE/DVT, pneumonia/atelectasis, an AMI, stroke or overdose / over-sedation could be the cause of the patient’s deterioration. Teachings advocated in the BTF Tier Two education (DETECT training) including A-G assessment apply when assessing patients with suspected sepsis.

Sepsis is a difficult clinical diagnosis that requires experience and a high index of suspicion for interpretation of history, signs and symptoms. Early senior clinician involvement is imperative to ensure that the required skills and knowledge are available to facilitate appropriate diagnosis and management. Suggested triggers for escalation to the AMO/infectious diseases physician are provided in the Toolkit Section B. The ISBAR framework can be used to structure the conversation to ensure completeness of information and standardize communication.

Sepsis treatment, reassessment and referral
The sepsis pathways provide clear resuscitation guidelines for patients with severe sepsis or septic shock sepsis using an A-G approach. The guideline is based on a bundle of care which should be delivered within one hour of recognition/diagnosis. The six components are:

1. Oxygen
2. Blood cultures
3. Serum lactate
4. Intravenous fluids
5. Antibiotics
6. Monitoring and reassessment
Reassessment of the patient is required to determine the effectiveness of treatment. Referral to a higher level of care such as an intensive care unit, specialist service or another facility may be required. The AMO should be informed that the patient has had a Clinical Review or Rapid Response call and commenced on the sepsis pathway.

Clinicians who suspect a paediatric patient of having sepsis should follow a stepped approach for escalation: Step 1 – facility (senior onsite medical officer), Step 2 – LHD (on-call paediatrician), Step 3 – State (NETS) as outlined in the NSW Health Policy: Recognition and Management of Patients who are Clinically Deteriorating (PD2013_049).

Communicating that the patient has been treated for sepsis and is at risk of further deterioration is essential at handover between shifts and when transferring patients to another ward/unit or hospital.

NOTE: The adult and paediatric sepsis pathways are not intended for patients at risk of febrile neutropenia. Patients with a recent haematological or oncology diagnosis should be managed using relevant local guidelines for febrile neutropenia.

Antibiotic guidelines
Prompt administration of antibiotics and resuscitation fluids is vital for effective management of sepsis in combination with effective source control. The goal is to commence antibiotic therapy within the first hour of recognition and diagnosis of sepsis. Antibiotic therapy should not be delayed whilst waiting for investigations or results.

The Inpatient Sepsis Intravenous Antibiotic Guideline aims to guide the prescription and timely administration of antibiotics for patients that have a diagnosis of sepsis and have been admitted to hospital for 48 hours or more. It is based on the recommendations in Therapeutic Guidelines: Antibiotic version 14, 2010 and is intended to provide an accessible resource which can be adapted to suit individual facility preferences in liaison with the antimicrobial stewardship team, local antimicrobial susceptibility patterns and senior clinicians.

Escalation to the Attending Medical Officer (AMO) for antibiotic advice
A flow chart is provided in the antibiotic guideline to assist the treating doctor in determining (for each case of sepsis) if the Antibiotic Guideline can be used or if escalation to the AMO for further advice is required.

Antibiotic prescribing for patients with sepsis due to hospital-acquired infections is complex because of:
- Unfamiliarity with the dosing, efficacy, toxicity, and spectrum of activity of rarely used and/or newer generation drugs e.g. meropenem, daptomycin, linezolid, tigecycline, colistin, amikacin, caspofungin
- Local resistance patterns of multi-drug resistant bacteria e.g. MRSA, VRE, ESBLs, MBLs, other MDR-GNBs and their association with different sources of infection
- Propensity for drug-resistance due to time spent in hospital, overseas travel, certain procedures, and previous antimicrobial therapy
- Constraints such as renal and/or hepatic failure, drug allergy and potential interactions with other medications
- Difficulty in assessing positive or negative microbiology results when rationalising therapy at 48-72 hours

Escalation to the infectious diseases physician or clinical microbiologist
In many cases, the AMO will feel comfortable prescribing antibiotics for such patients. It is anticipated however, that there will be times when the AMO recognises the triggers above as prompts for seeking immediate advice from a specialist Infectious Diseases Physician or Clinical Microbiologist.

A 24-hour a day referral pathway should exist for all NSW hospitals to obtain such advice on the authority of the patient’s AMO, who can then direct the junior medical officer (or registrar if possible) to call.
Review of antibiotic therapy
Antibiotic therapy must be reviewed within 24 hours of commencement and again when microbiology results are available. Antibiotics should be changed or ceased (if appropriate) when results are available. See the sepsis 48 hour management plan below.

Sepsis 48 hour management plan
Patients with a diagnosis of sepsis are at high risk of deterioration and require close monitoring and follow up care. The sepsis pathways are intended for the immediate management of sepsis only. Follow up management should be determined in consultation with the AMO and other expert clinicians.

A sepsis management plan is recommended to ensure that the patient is closely monitored, test results and antibiotic choices are reviewed in appropriate timeframes and the patient management plan is adjusted accordingly.

The CEC sepsis 48 hour management plan can be used for patients in the emergency department, ward or intensive care. The plan can be adapted to meet individual facility preferences.
EDUCATION

Resources have been developed that can be used by health care facilities to support widespread engagement with the program and develop clinician knowledge and skills. They can be adapted to meet health care facility needs and can be incorporated into existing education processes, such as BTF Tier Two education (DETECT training). It is highly recommended that local sepsis case studies and data are used.

The CEC provides a program of interactive webex / teleconference sepsis learning sessions. Details of the program and instructions for joining the sessions are distributed in advance via the LHD sepsis lead.

The following education resources are available on the CEC sepsis website and are periodically updated.

**SEPSIS KILLS video (4 minutes)**
Key messages on the **SEPSIS KILLS** program and why sepsis is a medical emergency. This video can be used as a stand-alone resource to quickly convey key messages or it can be incorporated into a wider education session.

**SEPSIS KILLS general presentation**
A presentation overview incorporating the five elements of the program and the NSW implementation strategy.

**SEPSIS KILLS clinical presentation: adults**
Key concepts of recognition, resuscitation and referral for adult patients with sepsis in the inpatient ward.

**SEPSIS KILLS clinical presentation: paediatrics**
Key concepts of recognition, resuscitation and referral for paediatric patients with sepsis in the inpatient ward.

**SEPSIS KILLS multiple choice questions**
This short quiz has evidence based questions to test sepsis knowledge and is suitable for a broad range of clinical staff caring for adult and/or paediatric patients.

**SEPSIS KILLS frequently asked questions**
The FAQ sheets address the commonly asked questions about the program.

**SEPSIS KILLS eLearning sepsis module**
This online module is available via the HETI Moodle site and includes adult and paediatric case studies on emergency department and ward patients.
EVALUATION

Evaluation is a systematic process to determine the extent of the improvements that have been made. Measurement ensures that clinical practice changes are carried out and provide a source of feedback and learning.

CEC sepsis database

The CEC sepsis database provides a tool to collect and analyse data. It is a web based application, which is accessed by entering hospital facility or LHD username and password on the NSW Health intranet system. CEC is working with HealthShare NSW to facilitate integration with other data collection programs.

The measures for the inpatient wards for both adults and paediatrics, have been agreed in consultation with clinicians and clinical governance units as follows:

<table>
<thead>
<tr>
<th>Measure</th>
<th>Description</th>
<th>Definition</th>
<th>Data collection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antibiotics</td>
<td>Timely administration of antibiotics increases the likelihood of survival and better patient outcomes</td>
<td>Time of CERS call / sepsis recognition to time of administration of first intravenous antibiotic</td>
<td>Time taken recorded on CEC sepsis data collection form that is then entered into CEC sepsis database</td>
</tr>
<tr>
<td>Time to administration of antibiotics</td>
<td>Timely administration of antibiotics increases the likelihood of survival and better patient outcomes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Resuscitation fluids</td>
<td>Timely administration of intravenous resuscitation fluids increases the likelihood of survival and better patient outcomes</td>
<td>Amount of bolus fluid administered within 60 minutes of CERS call / sepsis recognition</td>
<td>Record on CEC sepsis data collection form that is then entered into CEC sepsis database</td>
</tr>
<tr>
<td>Amount of bolus fluid administered</td>
<td>Timely administration of intravenous resuscitation fluids increases the likelihood of survival and better patient outcomes</td>
<td>Amount of bolus fluid administered within 60 minutes of CERS call / sepsis recognition</td>
<td>Record on CEC sepsis data collection form that is then entered into CEC sepsis database</td>
</tr>
<tr>
<td>Systolic Blood Pressure (Adults only)</td>
<td>Hypotension is a significant indicator in severe sepsis and septic shock</td>
<td>The first systolic blood pressure measurement taken at time of CERS call / sepsis recognition</td>
<td>SBP measurement level recorded on CEC sepsis data collection form that is then entered into CEC sepsis database</td>
</tr>
<tr>
<td>SBP recorded at time of CERS call / sepsis recognition</td>
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<tr>
<td>Clinical Review Call or Rapid Response</td>
<td>Indicates escalation of care is required</td>
<td>Level of escalation</td>
<td>Clinical Review Call or Rapid Response recorded on data collection form that is then entered into CEC sepsis database</td>
</tr>
<tr>
<td>Lactate</td>
<td>High serum lactate levels are an indicator of poor tissue perfusion and are strongly associated with morbidity and mortality</td>
<td>A serum lactate level collected within 60 minutes of CERS call / sepsis recognition</td>
<td>Collected Yes / No and if Yes serum lactate level recorded on CEC sepsis data collection form that is then entered into CEC sepsis database</td>
</tr>
<tr>
<td>Transfer of Care</td>
<td>Where the patient was managed or transferred to as a result of the sepsis episode</td>
<td>Ward, HDU / ICU, other hospital or tertiary referral</td>
<td>Ward, HDU / ICU, other hospital or tertiary referral recorded on CEC sepsis data collection form that is then entered into CEC sepsis database</td>
</tr>
</tbody>
</table>
Other measures that can be used to review care of patients with sepsis

Some additional measures that can be used at facility or LHD level to ascertain the effectiveness of the program include:

- Hospital length of stay for patients with sepsis
- Number of patients transferred to ICU with sepsis
- ICU length of stay for patients with sepsis
- Morbidity and mortality rates for patients with sepsis
- Root Cause Analysis data
- Clinical Review/Rapid Response data.

Monitoring and reporting your data

It is important to monitor, report and evaluate the program data to ensure that clinical practice and processes for recognition and management of the patient with sepsis are effective. The roles are outlined below.

**Facility**
Processes should be established for ward/unit and facility monitoring of progress. It is suggested that monitoring and report processes be integrated with existing Clinical Emergency Response Systems (CERS) for the deteriorating patient.

**Local Health District**
Clinical governance units have established monitoring and reporting requirements and systems at LHD level and will liaise with facilities on the local requirements.

**Clinical Excellence Commission**
Monitors and reports on statewide progress. Reports are provided six-monthly to LHD and facilities.
REFERENCES


13. Giaieski DF, Mikkelsen ME, Band RA, et al. Impact of time to antibiotics on survival in patients with severe sepsis or septic shock in whom early goal-directed therapy was initiated in the emergency department. Critical Care Medicine 2010; 38:1045-1053.

14. Australian Commission on Safety and Quality in Health Care, National Safety and Quality Health Service Standards. September 2012


GOVERNANCE

Program summary
Memo template
Facility implementation plan
Facility implementation checklist
Common causes of delay in recognising and treating sepsis in the wards
Implementation action plan
Ward implementation checklist
Sepsis issues log
SEPSIS KILLS PROGRAM

Sepsis is an international health care problem

Sepsis is a life-threatening condition that arises when the body’s response to infection injures its own tissues and organs, and can present in any patient in any clinical setting. It is one of the leading causes of death in hospital patients worldwide. More patients die from sepsis than prostate cancer, breast cancer and HIV/AIDS combined.

Severe sepsis and septic shock in adults have a mortality rate of around 25 per cent. In paediatric patients, sepsis is one of the leading causes of death with mortality rates as high as 10 per cent. Many of these deaths are preventable.

Appropriate recognition and timely management of patients with severe infection and sepsis is a significant problem in health care facilities. Delayed treatment is associated with high mortality rates, significant morbidity and high costs to the health care system. The mortality rate for adult patients with septic shock has been shown to increase by 7.6 per cent for every hour of delay in commencing antibiotic therapy, after the onset of hypotension. In the complex hospital ward environment there are frequently long delays between medical review and antibiotic prescription particularly when decision making is by junior medical officers.

SEPSIS KILLS program in NSW

The Clinical Excellence Commission (CEC) Clinical Focus Report on Recognition and Management of Sepsis found deficits in a range of clinical settings in NSW health care facilities. The lack of timely recognition and appropriate management was further demonstrated in the findings from the 2011 CEC Quality Systems Assessment.

The SEPSIS KILLS program aims to reduce preventable harm to patients through improved recognition and management of severe infection and sepsis in emergency departments and inpatient wards.

The program is based on three key actions:

- **RECOGNISE** the risk factors, signs and symptoms of sepsis
- **RESCUSITATE** with rapid intravenous fluids and administration of antibiotics within the first hour of diagnosis of sepsis
- **REFER** to appropriate senior clinicians and teams and retrieval if appropriate

It is founded on international evidence-based practice and brings significant benefits to NSW hospitals at both the clinical and system levels including:

- Enhanced clinician skills in sepsis recognition and management
- More timely, standardised and effective detection and management of sepsis
- Reduced mortality, morbidity and bed-stays from sepsis-related conditions
- Improved and safer patient experience.

The SEPSIS KILLS program was launched in May 2011 and has been implemented in 180 NSW public hospital emergency departments. The process of care for patients with sepsis has been greatly improved. Median time to start intravenous antibiotics has been reduced from more than 4 hours in 2011, to consistently less than 60 minutes in 2013 and 2014.

Inpatient Sepsis Program

Phase 2 of the SEPSIS KILLS program is focused on improving the recognition and management of sepsis in adults and children in inpatient wards of medium and large hospitals. It will be introduced Statewide from May 2014.
Local Health District has an Executive Sponsor and a sepsis lead. There will be an Executive-led sepsis team and clinical champions in each hospital driving the implementation and evaluation.

The inpatient program directly links with the Between the Flags (BTF) system and the NSW Health Policy: Recognition and Management of Patients who are Clinically Deteriorating (PD2013_049). Preliminary NSW sepsis data suggests that 30 per cent of clinical deterioration in adults that require a Rapid Response are septic. This is replicated in national and international literature with sepsis being a leading cause for clinical deterioration, accounting for one in three calls for a Rapid Response team.

A critical component of the sepsis program is timely and appropriate escalation to a senior clinician. The tools provided by the BTF system and SEPSIS KILLS program give clinicians objective criteria backed by policy for escalation, which empowers them to call for senior clinical help when they judge a patient to be deteriorating and/or are concerned that the patient has sepsis.

**Implementation in** (add hospital)

(Add local information here)

**Further information is available from:**

(Add facility/LHD sepsis leads here)

---

**Sepsis Program Lead**

Phone: (02) 9269 5500

Email: sepsis@cec.health.nsw.gov.au


**References**


INTERNAL MEMORANDUM

TO

FROM

TEL

DATE

SUBJECT SEPSIS KILLS program commencing at xxxx hospital

Sepsis is a medical emergency.

Sepsis is the cause of more adult deaths per year than prostate cancer, breast cancer and HIV combined. Thirty per cent of inpatients who require a Rapid Response are septic and mortality for these patients increases with delays in receiving appropriate treatment.

The SEPSIS KILLS program is going live in XX Hospital on xxxx 2014. It focuses on the Recognition, Resuscitation and Referral of patients who are septic.

Patients at greatest risk include those with:

- Any kind of infection-bacterial, viral, parasitic, or fungal-anywhere in the body
- Pre-existing (chronic) medical conditions
- Underactive immune system
- Surgery
- Invasive procedures or IV lines

When a patient is suspected of having sepsis, the medical officer must immediately undertake six key actions in discussion with a senior doctor:

1. Administer high flow oxygen to improve oxygen delivery to tissues
2. Take TWO blood cultures and other necessary specimens
3. Measure serum lactate - elevated levels are directly linked to increased mortality
4. Give appropriate IV antibiotics within 60 minutes. Every additional hour’s delay results in mortality increasing by 7.6%
5. Give adequate IV fluid resuscitation within 60 minutes to reduce organ dysfunction and multi-organ failure
6. Monitor urine output and vital signs after each fluid challenge and continue to re-assess

All xxxx are required to attend a session on the new sepsis recognition and management processes on xxxx 2014 and successfully complete the sepsis multiple choice questionnaire.
INSERT FACILITY NAME
IMPLEMENTATION PLAN
### Program Title:
SEPSIS KILLS program

### Program Aim:
To improve the recognition and timely management of sepsis in the inpatient wards of xxxx hospital

### Program Background:
*Include information here from the summary template provided in the Sepsis Toolkit plus local information to support the case for change*

### Program Benefits:
Implementation of the SEPSIS KILLS program will result in:
- enhanced clinician skills in sepsis recognition and management
- reduced mortality, morbidity and bed-stays from sepsis-related conditions
- enhanced networking opportunities across the system for clinicians and service teams
- improved quality and safety of care and a better and safer patient experience

### Program Objectives:
Use SMART objectives:
- **Specific**
- **Measurable**
- **Achievable**
- **Relevant**
- **Timely**
### SCOPE OF THE PROGRAM

<table>
<thead>
<tr>
<th>This program will include:</th>
<th>This program will not include:</th>
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<tbody>
<tr>
<td>Which clinical wards or units will be included, or will it be a whole of facility approach? Consider piloting in one or two wards or units before spreading to other areas.</td>
<td>What is out of scope?</td>
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</table>

#### Program Deliverables:

- What will you deliver at the end of the implementation process?
  
  NOTE: these are the products you will have at the end of the process, e.g. an education program, sepsis pathway adapted for local environments, improved awareness levels etc.

#### Program Milestones:

- Key activities and dates (month/year) they will be completed

#### Evaluation:

- How will you measure the success of the sepsis pathway implementation?
  
  NOTE: evaluation criteria must be specific and measurable eg
  
  - % clinical staff who attend an education session on sepsis recognition and management
  
  - % Rapid Responders who attend an education session on sepsis recognition and management and achieve 80% correct answers on the Sepsis MCQ
  
  - % of patients diagnosed with probable sepsis in xxxx facility who receive intravenous antibiotics within one hour of recognition of sepsis

#### Resources:

- What are the resources required to undertake the program? Consider: people, space to meet and access to a computer and internet, etc.

#### Linkages:

- Are there opportunities for this program to gain leverage or support from other groups? For example deteriorating patient program, national accreditation standards, clinical handover, antimicrobial stewardship, risk management programs.
### RISK ASSESSMENT

<table>
<thead>
<tr>
<th>Program Risks</th>
<th>Risk Rating</th>
<th>Mitigation Strategy</th>
<th>Residual Risk Rating</th>
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<tbody>
<tr>
<td>What are the risks to successful completion of the program?</td>
<td>(high, medium, low)</td>
<td>List strategies to remove or minimise the risks</td>
<td>(high, medium, low)</td>
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### COMMUNICATION PLAN

Who do you need to engage to make this program successful?

<table>
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<tr>
<th>Stakeholder</th>
<th>Position</th>
<th>What are their information needs?</th>
<th>How and when are you going to let them know?</th>
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# PROGRAM TEAM ROLES

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<th>Role</th>
<th>Details</th>
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| **Executive Sponsor:**| Name and designation of Executive Sponsor  
Role of the Executive Sponsor ie what do they do? |
| **Program Lead:**     | Name and designation  
Email  
Phone number  
Role of the Program Lead |
| **Clinical Lead(s):** | Name and designations  
Role of the Clinical Lead |
| **Program Team Members:** | Name and designations  
Role of the Program Team Members |
| **Start Date:**       | Completion Date: |

# ENDORSEMENT

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<th>Role</th>
<th>Details</th>
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</table>
| **Facility Executive Sponsor** | Name:  
Signature and Date: |
| **Facility Sepsis Lead** | Name:  
Signature and Date: |
| **LHD Sepsis Lead**     | Name:  
Signature and Date: |
| **LHD Director Clinical Governance** | Name:  
Signature and Date: |

## FACILITY IMPLEMENTATION CHECKLIST

### SEPSIS KILLS PROGRAM

#### GOVERNANCE

**Step 1: Establish a strong case for change**
- Develop a brief summary of the problem to capture interest.
- Specify the change to be made and provide evidence as to why the change should be made now.

**Step 2: Establish governance arrangements and a program team**
- Identify a program leader and executive sponsor
- Convene a team involving key stakeholders
- Identify the facility responsible committee and reporting/communication channels
- Develop a program plan for endorsement by hospital executive and LHD CGU
- Develop a comprehensive communication strategy.

**Step 3: Recruit respected and influential leaders and champions**
- Recruit medical and nursing leads/champions for each ward/unit
- Seek advice from emergency department sepsis clinical leads, Between the Flags coordinators and others who have led organisational clinical change programs.

**Step 4: Identify sepsis management barriers and enablers**
- Audit the baseline sepsis management evidence-practice gap
- Brainstorm the causes of inadequate or delayed recognition and treatment of sepsis
- Develop an action plan to address each barrier. Consider:
  - process change (WHAT has to be done); and
  - people change (HOW this will be communicated)
- Identify the enablers that will support the required changes.

#### CERS AND SEPSIS TOOLS

**Step 5: Adapt the sepsis tools to align with CERS and other local processes**
- Review the sepsis pathway and align with local CERS if needed
- Establish escalation process for patients with sepsis to:
  - AMO
  - Infectious Diseases
  - intensive care unit
- Develop/adapt local ISBAR tool for sepsis patients
- Establish minimum monitoring requirements for ward patients with sepsis
- Align the CEC sepsis antibiotic guideline with local drug and AMS processes

**Step 6: Identify tools and resources to support sepsis management**
- Consider use of sepsis blood order sets
- Identify components and placement of sepsis kit/pack/trolley
- Review IV cannulation access in-hours/after-hours
- Review the process for timely access to lactate measurement
### Step 7: Develop an education strategy
- Identify education tiers or groups to meet local needs
- Determine modes of education. Consider:
  - face to face awareness sessions
  - DETECT training
  - orientation
  - simulation training
- Document a plan to provide tailored education for specific professional and craft groups
- Determine education evaluation methodology.

### Step 8: Pilot the new sepsis process
- Implement the program in one or two areas using short Plan, Do, Study, Act cycles
- Consider use of an issues log to document concerns as they arise
- Seek feedback from stakeholders frequently and review progress
- Celebrate quick wins and adapt methods as needed
- Collect qualitative/quantitative data to measure improvement

### Step 9: Sustain and spread the approach to other wards/units
- Sustain the changes at the pilot site(s) and embed in organisational structures and routines
- Communicate the pilot outcomes using a variety of approaches including meetings, newsletters, intranet posts
- Market the good news stories and benefits to patients and staff
- Spread program to other wards/units in a phased approach
- Develop organisational guidelines on sepsis management principles and procedures

### Step 10: Measure, evaluate and improve
- Evaluate improvement – consider organisational, process and people change.
- Consider using a range of methods including:
  - medical record audit
  - observational audit
  - survey
  - focus group
  - consumer interviews
  - consumer feedback
- Display evaluation data and discuss at ward meetings
- Report results/action to staff, facility and LHD levels
- Promote success stories and lessons learnt.
COMMON CAUSES OF DELAY
RECOGNISING AND TREATING SEPSIS IN THE WARDS

Recognition
- Lack of nursing and junior doctor knowledge of sepsis risk factors, signs, symptoms
- Observations performed infrequently so not a timely indicator for deterioration and/or sepsis
- Sepsis screening processes different in each ward
- Rationalise that fever is disease related, not sepsis OR it can’t be sepsis as there is not fever
- Night RMO does not give high priority to sepsis due to workload and/or lack of knowledge that sepsis is a medical emergency
- Failure to communicate sepsis risk/diagnosis in clinical handover or ED to ward handover
- Not testing lactate to aid diagnosis due to lack of knowledge and/or access to testing
- Stuck in a particular diagnosis and sepsis not considered.

Escalation
- Senior clinician not available to assist in making diagnosis
- No formal RMO escalation process for sepsis
- Medical registrar, AMO or ID not consulted or informed of new sepsis
- Wait for investigations/specimens and/or results before escalation
- Surgical teams in theatre and unable to respond
- Long delay between call and after hours medical review
- NFR status unavailable
- Multiple teams looking after patient – confusion as to who to inform/seek advice.

Treatment
- Basic resuscitation not happening whilst awaiting arrival of Rapid Response team
- Doctor not familiar with sepsis pathway or bundle of care
- Long time to cannulation due to difficult access or not done at time of diagnosis
- Doctor or nurse cannulation skills not available on all shifts
- Fluid resuscitation volume/time for administration not standard
- Nurses (and doctors) don’t know what/how to give a rapid bolus (fluid challenge)
- Limited understanding of antibiotic prescribing choices/regimes in sepsis or give favoured antibiotics
- Senior doctor or Infectious Diseases advice not sought by junior doctors for initial treatment
- No antibiotics changed/commenced overnight – wait for morning rounds
- Wait for results of tests/investigations before commencing/changing antibiotics
- Prescription of antibiotics (writing up) and communication of same with nursing staff
- Nurses don’t consider antibiotics as urgent treatment
- Antimicrobial stewardship impacts on antibiotic choices and availability
- Antibiotics given as a slow infusion
- Equipment and resources for sepsis not centralised or available eg blood culture bottles
- Lack of monitoring of observations and urine output in patients post Rapid Response call or sepsis diagnosis.
## IMPLEMENTATION ACTION PLAN – SEPSIS KILLS PROGRAM

<table>
<thead>
<tr>
<th>Focus Area</th>
<th>Action</th>
<th>Owner</th>
<th>Due</th>
<th>Status / Progress</th>
</tr>
</thead>
<tbody>
<tr>
<td>Administration of antibiotics in 60 minutes or less</td>
<td>Implement process for MOs to check/establish IV access at the time of prescribing antibiotics.</td>
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<td></td>
<td>Set up sepsis trolley or packs in all wards with cannulation equipment and antibiotics.</td>
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<td></td>
<td>Review facility guidelines for administering antibiotics as IV bolus doses where appropriate.</td>
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## IMPLEMENTATION ACTION PLAN – SEPSIS KILLS PROGRAM

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</table>
## Ward Implementation Checklist

**Sepsis Kills Program**

<table>
<thead>
<tr>
<th>Governance</th>
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<tbody>
<tr>
<td>Nursing sepsis lead/champion identified</td>
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<tr>
<td>Medical sepsis lead/champion identified</td>
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<tr>
<td>Communication pathways for hospital sepsis lead and executive sponsor identified</td>
</tr>
<tr>
<td>SEPSIS KILLS is an agenda item for ward meetings</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CERS and Sepsis Tools</th>
</tr>
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<tbody>
<tr>
<td>Sepsis pathway displayed in ward areas</td>
</tr>
<tr>
<td>Paper and electronic versions of sepsis pathway readily available in nominated areas</td>
</tr>
<tr>
<td>Sepsis trolley or pack assembled with process agreed for checking and re-stocking</td>
</tr>
<tr>
<td>Sepsis blood order sets agreed</td>
</tr>
<tr>
<td>Antibiotic guideline and antibiotics available</td>
</tr>
<tr>
<td>Lactate measurement process agreed</td>
</tr>
<tr>
<td>IV cannulation available in-hours and after-hours</td>
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<tr>
<td>Escalation processes for nursing staff agreed</td>
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<tr>
<td>Escalation processes for medical staff agreed (AMO/senior clinician/ICU)</td>
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<tr>
<td>Minimum observation/monitoring requirements for patients with sepsis agreed</td>
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</table>

<table>
<thead>
<tr>
<th>Education</th>
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<tbody>
<tr>
<td>Program for SEPSIS KILLS education developed and communicated</td>
</tr>
<tr>
<td>Posters and other promotional materials available and displayed</td>
</tr>
<tr>
<td>All staff completed the SEPSIS KILLS awareness education</td>
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<tr>
<td>All staff completed the SEPSIS KILLS multiple choice questions</td>
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<tr>
<td>All staff educated on sepsis pathway data collection requirements</td>
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<table>
<thead>
<tr>
<th>Evaluation</th>
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<tbody>
<tr>
<td>Process for collection of sepsis pathways/audit sheets determined</td>
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<tr>
<td>Process for checking and collating sepsis data determined</td>
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<tr>
<td>SEPSIS KILLS results displayed in ward</td>
</tr>
<tr>
<td>Results/actions to improve discussed at ward meetings</td>
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<tr>
<td>Results/actions reported to sepsis lead/hospital responsible committee</td>
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<tr>
<td>Success stories and lessons learnt communicated in ward and other relevant forums</td>
</tr>
</tbody>
</table>
Please record all concerns and issues with the use and application of the Sepsis Pathway and antibiotic guideline.

Issues that should be logged include: difficulty with decision making or use of the pathway, escalation, communication and teamwork, medications and fluids, clinical incidents, and pathology.

All Clinical Incidents must be recorded in IIMS as per usual practice.

For support and advice please contact:

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<th>Name:</th>
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SEPSIS TOOLS

Adult inpatient sepsis pathway

Paediatric inpatient sepsis pathway

Sepsis 48 hour management plan

Adult inpatient sepsis intravenous antibiotic guideline

ISBAR tool

Escalation triggers for calling an AMO for patients with sepsis

Sepsis trolley/pack content list
SEPSIS PATHWAY – ADULT – INPATIENT

This pathway is intended for the recognition and immediate management of sepsis
Use relevant febrile neutropenia guidelines if the patient has haematology/oncology diagnosis

DOES YOUR PATIENT HAVE A KNOWN OR SUSPECTED INFECTION?

Does your patient have any of the following sepsis risk factors, signs or symptoms present?

- History of fevers or rigors
- Cough/sputum/breathlessness
- Abdominal pain/distension
- Line associated infection/redness/swelling/pain
- History of fevers or rigors
- Cough/sputum/breathlessness
- Abdominal pain/distension
- Line associated infection/redness/swelling/pain

Have a higher level of suspicion of sepsis for patients age > 65 years

PLUS

Does your patient have any RED ZONE observations or additional criteria?

No: LACTATE > 4mmol/L = Rapid Response

Yes

Patient has SEVERE SEPSIS or SEPTIC SHOCK until proven otherwise

- Sepsis is a medical emergency
- Call for a Rapid Response (as per local CERS) unless already made
- Commence treatment as per sepsis resuscitation guideline
- Inform the Attending Medical Officer that your patient has sepsis

Turn over page for sepsis resuscitation guideline

Patient may have SEPSIS

- Obtain senior clinician review
- Call for a Clinical Review (as per local CERS) unless already made
- Look for other causes of deterioration
- Commence treatment as per sepsis resuscitation guideline
- Inform the Attending Medical Officer (as per local CERS)

Turn over page for sepsis resuscitation guideline

Look for other common causes of deterioration

- New arrhythmia
- Hypovolaemia/haemorrhage
- Pulmonary embolus/DVT
- Atelectasis
- AMI
- Stroke
- Overdose/over sedation

- Initiate appropriate clinical care
- Repeat observations within 30 minutes AND increase the frequency of observations as indicated by the patient’s condition
- Re-evaluate for sepsis if observations remain abnormal or deteriorate

Discuss management plan with patient and family

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# SEPSIS PATHWAY – ADULT – INPATIENT

## SEPSIS RESUSCITATION GUIDELINE

<table>
<thead>
<tr>
<th><strong>A</strong></th>
<th><strong>B</strong></th>
<th><strong>C</strong></th>
<th><strong>D</strong></th>
<th><strong>E</strong></th>
<th><strong>F</strong></th>
<th><strong>G</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Maintain patent airway</td>
<td><strong>Give oxygen</strong>&lt;br&gt;Aim SpO₂ ≥ 95% (or 88-92% for COPD &amp; chronic type II respiratory failure)</td>
<td><strong>Large bore intravenous access, collect and check results:</strong>&lt;br&gt;☑ Lactate&lt;br&gt;☑ Blood cultures x 2&lt;br&gt;☑ FBC&lt;br&gt;☑ Blood gas&lt;br&gt;☑ EUC&lt;br&gt;☑ Procalcitonin if available&lt;br&gt;☑ Coags&lt;br&gt;☑ CRP&lt;br&gt;☑ LFTs&lt;br&gt;☑ Glucose</td>
<td><strong>Assess level of consciousness (LOC) using Alert, Verbal, Pain, Unresponsive (AVPU)</strong>&lt;br&gt;If V or less conduct a GCS&lt;br&gt;If P or U reassess Airway, Breathing and Circulation</td>
<td><strong>Examine patient for source of sepsis</strong>&lt;br&gt;Collect appropriate swabs, cultures, chest X-ray, ECG if indicated</td>
<td><strong>Fluid balance</strong>&lt;br&gt;Monitor and document fluid input &amp; output - consider IDC&lt;br&gt;Maintain urine output &gt; 0.5 mL/kg/hour</td>
<td><strong>Check Blood Glucose Level:</strong> if &gt; 12mmol/L consider glycaemic control</td>
</tr>
</tbody>
</table>

| **MONITOR & REASSESS** | **Continuously monitor and assess for signs of deterioration:**<br>• Respiratory rate in the Red or Yellow Zone<br>• SBP < 100mmHg<br>• Decreased or no improvement in level of consciousness<br>• Urine output < 0.5mL/kg/hour<br>• Increasing or no improvement in serum lactate |

<table>
<thead>
<tr>
<th><strong>REFER</strong></th>
<th><strong>THIS PATIENT HAS SEVERE SEPSIS OR SEPTIC SHOCK ESCALATION IN LEVEL OF CARE IS REQUIRED</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>This patient may need transfer to an Intensive Care Unit&lt;br&gt;• Discuss the patient’s condition with the Attending Medical Officer&lt;br&gt;• Consider a higher level of care as per local CERS&lt;br&gt;• Discuss management plan with patient and their family/carers</td>
<td><strong>Call for expert assistance after two failed IVC attempts</strong>&lt;br&gt;Escalate to Rapid Response if no response in SBP after 1000mL of fluid</td>
</tr>
</tbody>
</table>
SEPSIS PATHWAY – PAEDIATRIC – INPATIENT
This pathway is intended for the recognition and immediate management of sepsis
Use relevant febrile neutropenia guidelines if the patient has haematology/oncology diagnosis

ARE YOU CONCERNED THAT YOUR PATIENT COULD HAVE EARLY SEPSIS?

Does your patient have any of the following sepsis risk factors, signs or symptoms present?

- Deterioration despite treatment
- New or persistent signs of toxicity: Alertness, arousal or activity decreased; colour pale or mottled; cool peripheries; cry weak; grunting; rigors; bounding pulses; wide pulse pressure
- New onset of fever ≥38.5°C (neonates ≥38°C)
- High level parental concern
- Blood WCC outside normal range

AND

- Recent surgery cellulitis/wound infection
- Line associated infection/redness/swelling/pain
- Central line in-situ
- Chronic illness or immunocompromised
- 3 months of age or younger (corrected)

PLS

Does your patient have TWO or more YELLOW ZONE observations OR additional criteria OR clinician concern?

NB: Three Yellow Zone Criteria = Rapid Response

*Persistent tachycardia is consistent with sepsis

RECOGNISE

Does your patient have any RED ZONE observations OR additional criteria OR serious clinician concern?

Ensure senior clinician review to confirm likelihood of sepsis

RESPOND & ESCALATE

Patient has SEVERE SEPSIS or SEPTIC SHOCK until proven otherwise

- Sepsis is a medical emergency
- Call for a Rapid Response (as per local CERS) unless already made

Patient may have SEPSIS

- Obtain early senior clinician review within 30 minutes
- Consider and review blood gas

Any of these values are significant in sepsis

- Lactate ≥ 2 mmol/L
- BE ≤ - 5
- Procalcitonin (PCT) ≥ 0.5

Does senior clinician consider patient to be septic?

Commence treatment as per sepsis resuscitation guideline (over page)

Ensure the Attending Medical Officer/Paediatrician/NETS is aware (as per local CERS)

Discuss management plan with patient and family

Look for other common causes of deterioration

- Initiate appropriate clinical care
- Repeat observations within 30 minutes AND increase the frequency of observations as indicated by the patient’s condition
- Re-evaluate for sepsis if observations remain abnormal or deteriorate

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**SEPSIS PATHWAY – PAEDIATRIC – INPATIENT**

**SEPSIS RESUSCITATION GUIDELINE**

| **A** | Maintain patent airway |
| **B** | Give oxygen<br>Aim SpO₂ ≥ 95%<br>Monitor: Respiratory rate, Respiratory distress, SpO₂ |
| **C** | Intravenous access, collect and check results:<br> Blood gas<br> Lactate<br> EUC<br> Procalcitonin if available<br> Blood culture(s)<br> Coags<br> CRP<br> FBC<br> LFTs<br> Glucose<br> Consider intravenous access after two failed attempts or 60 seconds<br> IV Fluid Resuscitation<br> Give 20mL/kg 0.9% sodium chloride bolus STAT<br> Repeat 20mL/kg 0.9% sodium chloride bolus if no improvement in heart rate, capillary refill, colour<br> Monitor: Heart rate, BP, capillary refill, colour |
| **D** | Assess level of consciousness (LOC) using Alert, Verbal, Pain, Unresponsive (AVPU)<br> If V or less conduct a GCS<br> If P or U reassess Airway, Breathing and Circulation<br> Monitor: LOC |
| **E** | Examine patient for source of sepsis<br> Collect appropriate swabs, cultures, chest X-ray if indicated<br> Monitor: Temp |
| **F** | Fluid balance<br> Monitor and document fluid input & output - consider IDC<br> Maintain urine output ≥ 1mL/kg/hour<br> Monitor: Urine output |
| **G** | Check Blood Glucose Level:<br> if less than 3mmol/L treat with 2mL/kg 10% Dextrose<br> Monitor: BGL |

**PRESCRIBE and ADMINISTER ANTIBIOTICS WITHIN 60 MINUTES**

**Do not delay for investigations or results**

If patient already on antibiotic therapy this **MUST** be reviewed by the Attending Medical Officer<br> Consider alternate source of infection and/or resistance

**MONITOR & REASSESS**

**Continue monitoring and assess for signs of deterioration:**
- Persistent tachycardia, slow capillary refill and hypotension
- Colour pale and mottled
- Drowsiness or abnormal LOC
- Urine output < 1mL/kg/hour
- Acidosis, increasing serum lactate or procalcitonin
- Hypoglycaemia, leukopenia or abnormal coagulation

**THIS PATIENT HAS SEVERE SEPSIS OR SEPTIC SHOCK**

**ESCALATION IN LEVEL OF CARE IS REQUIRED**

This patient may need transfer to a Paediatric Intensive Care Unit
- Update the Attending Medical Officer on patient’s condition
- Seek advice from local/regional paediatric experts and/or NETS Tel: 1300 36 2500
- Early inotropes and intubation may be necessary
- Discuss management plan with patient and their family/carers

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## SEPSIS 48 HOUR MANAGEMENT PLAN

<table>
<thead>
<tr>
<th>Time</th>
<th>Action</th>
<th>Criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>0 – 2 Hours</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Communication</td>
<td>Attending Medical Officer (AMO) informed that patient is on sepsis pathway</td>
<td>Clinical handover <strong>must</strong> inform the receiving team that the patient was treated for sepsis</td>
</tr>
</tbody>
</table>
| Monitor and reassess | Monitor and reassess for sepsis deterioration which may include one or more of the following: | - *Respiratory rate in the Red or Yellow Zone*  
  - *Systolic blood pressure < 100mmHg*  
  - *Decreased or no improvement in level of consciousness*  
  - *Urine output less than 0.5mL/kg/hr*  
  - *No improvement in serum lactate level*  
  
  If deteriorating, activate local CERS and inform AMO |
| Sepsis screen | Head to toe assessment for infection source and initiate investigations which may include: | - Diagnostic imaging  
  - Urine MSU (or CSU) for MCS  
  - Sputum for MCS  
  - Faeces for C.difficile and MCS (if diarrhoea)  
  - Wound swab for MCS  
  - Nasopharyngeal swabs  
  - Lumbar puncture (if indicated) |
| Antibiotics | Medical officer to consult with AMO on appropriate antibiotic prescribing | | Prescribe antibiotics in the medication chart and indicate the appropriate time for dosing |
| IV Fluids | Prescribe IV fluids as appropriate  
  Monitor haemodynamic observations | | |

---

**Recognition - Resuscitate - Refer**

[Image: Sepsis kills logo]
<table>
<thead>
<tr>
<th>Time</th>
<th>Action</th>
<th>Criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>2 – 24 Hours</strong></td>
<td><strong>Continue monitoring</strong></td>
<td>Monitor and reassess for sepsis deterioration which may include one or more of the following:</td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Respiratory rate in the Red or Yellow Zone</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Systolic blood pressure &lt; 100mmHg</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Decreased or no improvement in level of consciousness</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>Urine output less than 0.5mL/kg/hr</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td><em>No improvement in serum lactate level</em></td>
</tr>
<tr>
<td></td>
<td></td>
<td>If deteriorating, activate local CERS and inform AMO</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If improving, continue observations every 30 minutes for 2 hours, then hourly for 4 hours</td>
</tr>
<tr>
<td><strong>Repeat lactate</strong></td>
<td></td>
<td>Lactate level 4 hours post recognition</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Date -- /--/ ---- Time --:-- mmol/L</td>
</tr>
<tr>
<td><strong>Fluid Resuscitation</strong></td>
<td></td>
<td>Prescribe IV fluids as appropriate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Monitor haemodynamic observations</td>
</tr>
<tr>
<td><strong>Reassess</strong></td>
<td></td>
<td>Check preliminary blood counts</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If patient is neutropenic, review antibiotics and change if needed</td>
</tr>
<tr>
<td><strong>Review treatment</strong></td>
<td></td>
<td>Review need for advanced care directives and ceiling of care if appropriate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Are any other causes of deterioration likely?</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Discuss with AMO and treat accordingly</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Cease antibiotics if appropriate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Continue monitoring for deterioration including urine output</td>
</tr>
<tr>
<td><strong>24 – 48 Hours</strong></td>
<td><strong>Reassess</strong></td>
<td>Repeat biochemistry as indicated</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Review results of tests and investigations</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Discuss with AMO and treat accordingly</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Cease antibiotics if appropriate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Continue monitoring for deterioration including urine output</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Confirm diagnosis and document source of sepsis in medical record</td>
</tr>
</tbody>
</table>

This document is intended for patients who have been recognised as having sepsis and have started on a sepsis pathway. The Sepsis 48 Hour Management Plan aims to guide clinical staff using a step by step process which ensures that the patient monitoring and treatment is appropriate.
The Clinical Excellence Commission (CEC) Adult Inpatient Sepsis Intravenous Antibiotic Guideline aims to guide the prescription and timely administration of antibiotics for adult inpatients that have a diagnosis of sepsis, severe sepsis or septic shock and have been admitted to hospital for 48 hours or more.

The guideline is based on the recommendations in Therapeutic Guidelines: Antibiotic version 14, 2010. It is intended to provide an accessible resource, which can be adapted to suit individual facility preferences in liaison with the antimicrobial stewardship team and local antimicrobial susceptibility patterns. Antimicrobial stewardship teams may wish to refer to their latest hospital cumulative antibiogram, if available, when modifying the guideline.

Prompt administration of antibiotics and resuscitation fluids is vital in the management of the patient with sepsis. The goal is to commence antibiotic therapy within the first hour of the recognition and diagnosis of severe sepsis. The selection of appropriate antimicrobial therapy is complex and this guideline is not intended to cover all possible scenarios.

Clinicians must review antimicrobial therapy within 24 hours of commencement, and change or cease antibiotics as required once microbiology results are available.

This guideline is not intended for:

- patients with FEBRILE NEUTROPENIA who should be managed using local febrile neutropenia guidelines
- small hospitals and multi-purpose services where it would be more appropriate to use the Sepsis Adult FIRST DOSE Empirical Intravenous Antibiotic Guideline – Emergency Department
- patients who are deemed to have had incubating or unrecognised community acquired sepsis on admission. Use the Sepsis Adult FIRST DOSE Empirical Intravenous Antibiotic Guideline – Emergency Department

Obtain at least two sets of blood cultures from separate venepuncture sites before antibiotic administration.

Obtain other clinical specimens as appropriate but do not delay administration of antibiotics or wait for results of investigations.

The antimicrobial treatment indication and plan should be documented in the patient record.
Patient meets sepsis pathway criteria

Does the patient have one or more of the following?

- antibiotic therapy within the last 7 days
- had a recent infection with a multi-resistant organism (MRO)* or is known to be colonised with an MRO
- contra-indications to specific antimicrobial therapy recommended in the guideline
- multiple possible sources of infection
- acute renal and/or hepatic failure
- risk factors for an antibiotic resistant infection due to time spent in hospital(s), overseas hospitalisation or residential care in previous 12 months
- surgical procedures that may influence the likely source of infection (e.g. urological surgery).

Yes

Does the patient have febrile neutropenia?

No

Consult immediately with the Attending Medical Officer regarding antibiotic choice. Advice from the designated infectious diseases and/or clinical microbiology services may be required

Yes

Consult immediately the Attending Medical Officer and manage according to febrile neutropenia guideline relevant for your facility

No

Follow antibiotic regimen outlined in Table 1 or local guideline if available in your facility

---

*Examples of MROs include methicillin-resistant Staphylococcus aureus (MRSA), vancomycin-resistant enterococci (VRE), extended-spectrum beta-lactamase (ESBL) producing organisms and carbapenem-resistant Gram negative organisms.
<table>
<thead>
<tr>
<th>Apparent source of sepsis</th>
<th>Sepsis antibiotic regimen</th>
<th>Penicillin allergic not immediate hypersensitivity</th>
<th>Penicillin or cephalosporin allergic immediate hypersensitivity or severe prior reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sepsis secondary to hospital acquired pneumonia, low risk of MRO (generally patient who has been in hospital &lt; 5 days who does not have risk factors for MRO)</td>
<td>ceftriaxone 1 g IV, daily</td>
<td>ceftriaxone 1 g IV, daily</td>
<td>moxifloxacin 400 mg IV, daily</td>
</tr>
<tr>
<td></td>
<td>OR</td>
<td>OR</td>
<td></td>
</tr>
<tr>
<td></td>
<td>benzylpenicillin 1.2 g IV, 6-hourly PLUS</td>
<td>cefotaxime 1 g IV, 8-hourly</td>
<td></td>
</tr>
<tr>
<td></td>
<td>gentamicin 4 to 6 mg/kg IV, for 1 dose (severe sepsis 7 mg/kg)</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>OR</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>cefotaxime 1 g IV, 8-hourly</td>
<td></td>
<td></td>
</tr>
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<td></td>
<td>OR</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>piperacillin+tazobactam 4+0.5 g IV, 8-hourly</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>OR</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ticarcillin+clavulanate 3+0.1 g IV, 6-hourly</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sepsis secondary to hospital acquired pneumonia, high risk of MRO</td>
<td>piperacillin+tazobactam 4+0.5 g IV, 6-hourly</td>
<td>cefepime 2 g IV, 8-hourly</td>
<td>Seek expert advice</td>
</tr>
<tr>
<td></td>
<td>OR</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ticarcillin+clavulanate 3+0.1 g IV, 6-hourly</td>
<td></td>
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<td></td>
<td>OR</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>cefepime 2 g IV, 8-hourly</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>If the patient is ventilated ADD</td>
<td></td>
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<tr>
<td></td>
<td>gentamicin 4 to 6 mg/kg IV, for 1 dose (severe sepsis 7 mg/kg)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>If MRSA prevalent in your hospital</td>
<td>ADD</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>vancomycin 1.5g IV, 12-hourly</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apparent source of sepsis</td>
<td>Sepsis antibiotic regimen</td>
<td>Penicillin allergic not immediate hypersensitivity</td>
<td>Penicillin or cephalosporin allergic Immediate hypersensitivity or severe prior reaction</td>
</tr>
<tr>
<td>----------------------------------------------------------------</td>
<td>------------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Severe sepsis with an apparent urinary tract source</td>
<td>gentamicin 4-7 mg/kg IV, for 1 dose PLUS ampicillin 2 g IV, 6-hourly</td>
<td>gentamicin 4-7 mg/kg IV, for 1 dose</td>
<td>gentamicin 4-7 mg/kg IV, for 1 dose</td>
</tr>
<tr>
<td></td>
<td></td>
<td>OR</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>ceftriaxone 1 g IV, daily if gentamicin is contraindicated</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>OR</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>cefotaxime 1 g IV, 8-hourly if gentamicin is contraindicated</td>
<td></td>
</tr>
<tr>
<td>Severe sepsis with an apparent biliary or gastrointestinal tract source</td>
<td>ampicillin 1 g IV, 6-hourly PLUS gentamicin 4 to 7 mg/kg IV, for 1 dose PLUS metronidazole 500 mg IV, 12-hourly</td>
<td>metronidazole 500 mg IV, 12-hourly PLUS ceftriaxone 1 g IV, daily</td>
<td>gentamicin 4 to 7 mg/kg IV, for 1 dose AND seek expert advice</td>
</tr>
<tr>
<td></td>
<td></td>
<td>OR</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>metronidazole 500 mg IV, 12-hourly PLUS cefotaxime 1 g IV, 8-hourly</td>
<td></td>
</tr>
<tr>
<td>Severe sepsis resulting from a skin infection (including cellulitis) or surgical site infection</td>
<td>flucloxacillin 2 g IV, 6-hourly</td>
<td>cephalozin 2 g IV, 8-hourly</td>
<td>clindamycin 450 mg IV, 8-hourly</td>
</tr>
<tr>
<td></td>
<td>If MRSA prevalent in your hospital ADD vancomycin 1.5g IV, 12-hourly</td>
<td>If MRSA prevalent in your hospital ADD vancomycin 1.5g IV, 12-hourly</td>
<td>OR</td>
</tr>
<tr>
<td></td>
<td></td>
<td>OR</td>
<td></td>
</tr>
<tr>
<td></td>
<td>If likely to be MRSA colonized ADD vancomycin 1.5g IV, 12-hourly</td>
<td>If patient meets criteria for toxic shock ADD clindamycin 600mg IV, 8-hourly</td>
<td>Seek expert advice</td>
</tr>
<tr>
<td>Maternal sepsis (peri or post-partum) if source unclear</td>
<td>piperacillin+tazobactam 4+0.5 g IV, 8-hourly</td>
<td>ceftriaxone 1g IV, 24-hourly PLUS metronidazole 500mg IV 12-hourly</td>
<td></td>
</tr>
<tr>
<td></td>
<td>If patient meets criteria for toxic shock ADD clindamycin 600mg IV, 8-hourly</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>If likely to be MRSA colonized ADD vancomycin 1.5g IV, 12-hourly</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>If patient meets criteria for toxic shock ceftriaxone 1 g IV, 24-hourly PLUS clindamycin 600mg IV, 8-hourly</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**TABLE 1: ANTIBIOTIC PRESCRIBING (review after 24 hours)**
### TABLE 1: ANTIBIOTIC PRESCRIBING (review after 24 hours)

<table>
<thead>
<tr>
<th>Apparent source of sepsis</th>
<th>Sepsis antibiotic regimen</th>
<th>Penicillin allergic not immediate hypersensitivity</th>
<th>Penicillin or cephalosporin allergic Immediate hypersensitivity or severe prior reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maternal sepsis likely to be due to Group A streptococcal infection</td>
<td>benzylpenicillin 2.4g IV, 4-hourly PLUS clindamycin 600mg iv 8-hourly</td>
<td>cephalosolin 2 g IV, 6-hourly PLUS clindamycin 600mg iv 8-hourly</td>
<td>Seek expert advice</td>
</tr>
<tr>
<td>OR</td>
<td>benzylpenicillin 2.4g IV, 4-hourly PLUS lincomycin 600 mg IV 8-hourly</td>
<td>cephalosolin 2 g IV, 6-hourly PLUS lincomycin 600 mg IV 8-hourly</td>
<td></td>
</tr>
<tr>
<td>Severe sepsis, unknown source or focus, including possible IV line-associated sepsis Removal of the infected IV device is usually required</td>
<td>flucloxacinillin 2 g IV, 6-hourly PLUS gentamicin 4-7 mg/kg IV, for 1 dose</td>
<td>cephalosolin 2 g IV, 8-hourly PLUS gentamicin 4-7 mg/kg IV, for 1 dose</td>
<td>vancomycin 1.5 g IV, 12-hourly PLUS gentamicin 4-7 mg/kg IV, for 1 dose</td>
</tr>
<tr>
<td>If MRSA prevalent in your hospital ADD vancomycin 1.5g 12-hourly</td>
<td>If MRSA prevalent in your hospital ADD vancomycin 1.5g 12-hourly</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### NOTES FOR TABLE 1

- **Definitions of penicillin hypersensitivity**
  - **Immediate hypersensitivity** involves the development of urticaria, angioedema, bronchospasm or anaphylaxis within one to two hours of drug administration.
  - **Severe prior reaction** involves a history of drug rash eosinophilia and systemic symptoms (DRESS) or Stevens-Johnson Syndrome following administration of a penicillin or cephalosporin.
  - All penicillin and cephalosporin class antibiotics are contraindicated in patients with history of drug rash eosinophilia and systemic symptoms (DRESS), Stevens-Johnson Syndrome or IgE-mediated immediate penicillin or cephalosporin allergy.

- **Definitions of low risk and high risk of MRO**

- **Doses for renal impairment (creatinine clearance ≤ 60mL/min)**
| Gentamicin and vancomycin dosing and frequency | Refer to *Therapeutic Guidelines: Antibiotic* for more information  
|-----------------|--------------------------------------------------|
| Criteria for toxic shock | Refer to *Therapeutic Guidelines: Antibiotic* for more information  
| Notes for gentamicin | One dose of gentamicin is recommended; for subsequent doses, assess renal function and adjust frequency accordingly  
Use for a maximum of 48 hours as empirical therapy pending outcome of investigations; monitoring of plasma concentrations NOT required if gentamicin is not used beyond 48 hours  
Directed therapy (beyond 48 hours, based on microbiology results) should be used on the advice of infectious diseases physician or clinical microbiologist only  
Dose should be based on ideal body weight or actual body weight – *whichever of the two is lower*  
The maximum dose of gentamicin in severe sepsis is 640 mg  
For other indications, the maximum dose is lower. Refer to Table 2.24 in *Therapeutic guidelines: Antibiotic*, version 14, 2010.  
**Contraindications:**  
Previous vestibular or auditory toxicity due to an aminoglycoside  
Serious hypersensitivity reaction to an aminoglycoside  
**Precautions:**  
Pre-existing significant hearing problems  
Pre-existing vestibular problems  
Neuromuscular disorders, including myasthenia gravis  
Chronic liver disease or severe cholestasis (bilirubin above 90 micromol/L)  
Chronic renal failure or deteriorating renal function – consult AMO |
Reconstitute antibiotics with sterile water for injection (WFI) unless stated otherwise.

If further dilution is required for IV injection or infusion, use sterile sodium chloride 0.9% or sterile glucose 5% unless stated otherwise.

Where possible use separate dedicated lines for resuscitation fluid and for medications. When injecting antibiotics directly into an IV injection port which has resuscitation fluid running:
- clamp the infusion fluid line and flush with 20 mL sterile sodium chloride 0.9% solution
- administer antibiotic over the required time
- flush the line with 20 mL sterile sodium chloride 0.9% solution and recommence resuscitation fluid.

<table>
<thead>
<tr>
<th>Antibiotic</th>
<th>Presentation (adult)</th>
<th>Reconstitution fluid/volume</th>
<th>Final volume</th>
<th>Minimum administration time</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>ampicillin</td>
<td>Vial 1 g</td>
<td>10 mL WFI</td>
<td>10 - 20 mL</td>
<td>3 – 5 minutes</td>
<td>Penicillin class antibiotic</td>
</tr>
<tr>
<td>benzyl/penicillin</td>
<td>Vial 600 mg</td>
<td>2 mL WFI</td>
<td>10 mL</td>
<td>3 – 5 minutes</td>
<td>Penicillin class antibiotic</td>
</tr>
<tr>
<td></td>
<td>Vial 1.2 g</td>
<td>4 mL WFI</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>cefepime</td>
<td>Vial 1 g</td>
<td>10 mL NS</td>
<td>10 mL</td>
<td>3 - 5 minutes</td>
<td>Cephalosporin class antibiotic</td>
</tr>
<tr>
<td>ceftriaxone</td>
<td>Vial 1 g</td>
<td>10 mL WFI</td>
<td>10 mL</td>
<td>2 – 4 minutes</td>
<td>Cephalosporin class antibiotic incompatible with calcium containing solutions, flush thoroughly before and after with sodium chloride 0.9%</td>
</tr>
<tr>
<td>cefotaxime</td>
<td>Vial 1 g</td>
<td>10mL WFI</td>
<td>10 mL</td>
<td>3 – 5 minutes</td>
<td>Cephalosporin class antibiotic</td>
</tr>
<tr>
<td>cephazolin</td>
<td>Vial 1 g</td>
<td>10 mL WFI</td>
<td>10 mL</td>
<td>3 – 5 minutes</td>
<td>Cephalosporin class antibiotic</td>
</tr>
<tr>
<td>clindamycin</td>
<td>Ampoules 300 mg/2 mL</td>
<td>N/A</td>
<td>600 mg in 50 mL</td>
<td>20 minutes</td>
<td>Check product is clear of any crystals prior to administration</td>
</tr>
<tr>
<td></td>
<td>600 mg/4 mL</td>
<td></td>
<td>900 mg in 100 mL</td>
<td>30 minutes</td>
<td></td>
</tr>
<tr>
<td>Antibiotic</td>
<td>Presentation (adult)</td>
<td>Reconstitution fluid/volume</td>
<td>Final volume</td>
<td>Minimum administration time</td>
<td>Notes</td>
</tr>
<tr>
<td>------------------------------</td>
<td>----------------------</td>
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<td>----------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| flucloxacillin               | Vial 1 g             | 5 mL WFI                    | 10 mL        | 3 - 5 minutes (1 g)          | Penicillin class antibiotic  
Repeated doses of 2 g via a peripheral line should be further diluted and infused over 20 – 30 minutes|
|                              |                      |                            | 20 mL        | 10 minutes (2 g)             |                                                                                                                                         |
| gentamicin                   | Ampoule 80 mg/2 mL   | N/A                        | 10-20 mL     | (240mg or less) 3 – 5 minutes | Refer to notes for gentamicin                                                                                                             |
|                              |                      |                            |              |                             | 50 mL or 100 mL (more than 240mg) 30 minutes                                                                                          |
| lincomycin                   | Vial 600mg/2mL       | N/A                        | 100mL        | 60 minutes                  |                                                                                                                                         |
| metronidazole                | Infusion bag 500 mg /100 mL | N/A                       | See presentation column | 20 minutes                  |                                                                                                                                         |
| moxifloxacin                 | Infusion bag 400 mg /250 mL | N/A                       | See presentation column | 60 minutes                  | May prolong QT interval and lead to ventricular arrhythmias. May induce seizures in epileptics                                      |
| piperacillin with tazobactam | Vial 4 g/0.5 g       | 20 mL WFI                   | 50 mL        | 30 minutes                  | Penicillin class antibiotic                                                                                                           |
| ticarcillin with clavulanic acid | Vial 3 g/0.1 g   | 13 mL WFI                   | 50 mL        | 30 minutes                  | Penicillin class antibiotic                                                                                                           |
| vancomycin                   | Vial 500 mg          | 10 mL WFI                   | Dilute to maximum concentration of 5mg/mL for peripheral line | Maximum of 10 mg/minute       | Infusion related effects are common, decrease infusion rate and monitor closely if these occur                                           |
|                              | Vial 1 g             | 20 mL WFI                   |              |                             |                                                                                                                                         |
FURTHER MANAGEMENT

The patient should be reviewed by the Attending Medical Officer within 24 hours of commencing the sepsis pathway and antibiotic therapy, with referral to the infectious diseases and/or clinical microbiology services for specific advice.

Microbiology results are generally available within 48-72 hours, and should be used to guide further management of the patient. This may include de-escalating or ceasing antimicrobial therapy.

Clinicians that are experiencing difficulty in assessing positive or negative microbiology results when rationalising antibiotic therapy at 48-72 hours should contact the designated infectious diseases and/or clinical microbiology services.

Check local instructions regarding referral

References


# ISBAR TOOL

## Introduction

State:
- Your name
- Your role/position in the hospital
- The ward/unit and hospital/health service you are calling from
- I am calling about my patient.....identify the patient (e.g. name, MRN, DOB)

NB: Patients 65+ years of age are at greater risk of sepsis

## Situation

I am concerned they are deteriorating because...
- Detail the clinical problem that is prompting the call - I think this patient is septic
- The patient is stable/unstable

## Background

- Symptoms - state if symptoms meets the sepsis pathway
- Vital signs including any trends
- Medical history pertinent to the reason for calling
  - Patients who are immune compromised and have a chronic illness are at greater risk of sepsis
  - If recently taken or currently on antibiotics
- Any initial treatment you have commenced, including whether or not it has worked. Sepsis bundle commenced oxygen, blood cultures, lactate, IV fluids, antibiotics (if given)

## Assessment

- What do you think the problem is?
- Are there any risks you have identified? Any risk factors of sepsis
- Clinical signs that support your diagnosis or impression. State if symptoms meets the sepsis pathway.
- Any tests or procedures you think need to be done and blood results including lactate.
- Any test(s)/scan(s) results you are waiting for

## Recommendation

Be clear about what you would like to happen, what you want the person you are calling to do and who is responsible for these actions e.g. Urgent review, transfer to another facility or advise on treatment
- I need you to come to review the patient
- I need you to authorise transfer of the patient
- I need you to authorise the test/scan
- I need you to follow up on
- I would like some guidance on
  - further management of the patient
  - antibiotic prescribing
  - further tests or considerations
- I think that this patient would be better managed in a higher level of care
# ESCALATION TRIGGERS

## FOR CALLING AN AMO FOR PATIENTS WITH SEPSIS

<table>
<thead>
<tr>
<th>Escalation triggers</th>
<th>Considerations for escalation to ID / micro specialist</th>
</tr>
</thead>
</table>
| ▪ Rapid Response call  
  - including serum lactate > 4mmol/L (adults)  
  - serum lactate > 2mmol/L is a concern in paediatric patients  
| ▪ 3 Clinical Review Calls in an eight hour timeframe  
| ▪ 3 simultaneous Yellow Zone criteria in a paediatric patient  
| ▪ Persisting hypotension (SBP less than 90mmHg)  
| ▪ High level of clinician concern  
| ▪ Serious concern by any patient/family member  
| ▪ Significant deterioration in condition and/or admission to ICU  
| ▪ Patient is immunocompromised  
| ▪ Patient with complex antibiotic prescribing requirements  |
| ▪ Unfamiliarity with the dosing, efficacy, toxicity, and spectrum of activity of rarely used and/or newer generation drugs (e.g., meropenem, daptomycin, linezolid, tigecycline, colistin, amikacin, caspofungin, cefepime)  
| ▪ Patient has had a recent infection with a resistant organism  
| ▪ Pre-existing antibiotics, recent or current antibiotic therapy  
| ▪ Any contradictions to specific antimicrobial therapy  
| ▪ Multiple possible sources of infection  
| ▪ Acute renal failure  
| ▪ Recent travel overseas or to a tropical location  
| ▪ Difficulty in assessing positive or negative microbiology results when rationalising therapy at 48-72 hours.  
| ▪ Recent candidiasis  |

The triggers and considerations are suggested as a guide for staff, and can be adapted to suit local facility and LHD requirements.
<table>
<thead>
<tr>
<th>Item</th>
<th>Detail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sepsis pathway</td>
<td></td>
</tr>
<tr>
<td>Antibiotic guideline</td>
<td></td>
</tr>
<tr>
<td>Sepsis 48 hour guideline</td>
<td></td>
</tr>
<tr>
<td>Data collection form</td>
<td></td>
</tr>
<tr>
<td>Personal protective equipment</td>
<td></td>
</tr>
<tr>
<td>Venepuncture / cannulation equipment</td>
<td></td>
</tr>
<tr>
<td>Antibiotic vials as per local guidelines</td>
<td></td>
</tr>
<tr>
<td>IV resuscitation fluid</td>
<td></td>
</tr>
<tr>
<td>Giving sets (including blood pump set)</td>
<td></td>
</tr>
<tr>
<td>Blood culture bottles</td>
<td></td>
</tr>
<tr>
<td>Pathology tubes</td>
<td></td>
</tr>
<tr>
<td>Pathology forms</td>
<td></td>
</tr>
<tr>
<td>Syringes</td>
<td></td>
</tr>
</tbody>
</table>
EDUCATION

- Sepsis video (available from CEC website)
- Sepsis program general presentation (available from CEC website)
- Sepsis clinical presentation – adults (available from CEC website)
- Sepsis clinical presentation – paediatrics (available from CEC website)
- Sepsis multiple choice questions (Toolkit and available from CEC website)
- Sepsis multiple choice answers (Toolkit and available from CEC website)
SEPSIS KILLS PROGRAM
MULTIPLE CHOICE QUESTIONS

Q1. What is sepsis?

A) Any infection
B) An infection plus hypotension
C) An infection plus systemic inflammatory response syndrome
D) Someone who is on antibiotics

Q2. Which signs and symptoms are most common in adult patients with severe sepsis?

A) Fever > 38deg C
B) Tachycardia
C) Tachypnoea
D) Metabolic acidosis
E) Acute oliguria

Q3. Which of the following criteria present and new in your patient should prompt commencement of the Sepsis Pathway?

A) RR - 20 breaths/min
B) RR - 20 breaths/min + HR > 122bpm
C) RR > 25 breaths/min + HR > 120bpm + recent surgery

Q4. Patients with severe sepsis and a lactate level $\geq$ 4 mmol/L have an increased likelihood of mortality compared to patients with a normal lactate.

A) True
B) False

Q5. In patients with sepsis, which of the following statements is incorrect?

A) Antibiotics should never be delayed while waiting for the results of investigations.
B) Patients with febrile neutropenia should be managed using the local febrile neutropenia guideline
C) Time to first dose antibiotics does not influence outcome
D) The patient should be reviewed by the attending medical officer within 24 hours of starting on sepsis pathway and antibiotic therapy, with referral to the infectious diseases / clinical microbiology service for specific advice.
Q6. Which signs and symptoms are most common in children and infants with sepsis?

A) Fever >38deg C  
B) Persistent Tachycardia  
C) Lethargy  
D) Metabolic acidosis  
E) Lactate >4mmol/L

Q7. During the monitor and reassess phase of resuscitation of the septic child or infant, which of the following signs of deterioration indicates your patient requires escalation in level of care?

A) Persistent tachycardia, slow capillary refill and hypotension  
B) Drowsiness or abnormal LOC and/or urine output<1ml/kg/hr  
C) Acidosis, increasing serum lactate or procalcitonin  
D) Hypoglycaemia, leukopenia or abnormal coagulation  
E) Any of the above

Q8. BP is a reliable parameter in the early recognition of sepsis in paediatric patients

A) True  
B) False

Q9. Fluid resuscitation in paediatric sepsis should be given as:

A) 10ml/kg 0.9% sodium chloride over 20minutes  
B) 20ml/kg bolus of 0.9% sodium chloride over no more than 10minutes  
C) 20ml/kg bolus of 0.9% sodium chloride over 60 minutes  
D) 10ml/kg 0.9% sodium chloride over 10minutes

Q10. Key messages are: SEPSIS KILLS & TIME IS LIFE. Fill in the boxes:

- risk factors, signs, symptoms of sepsis and have early involvement of senior clinicians
- with rapid IV fluids)and administer antibiotics within one hour
- to appropriate in hospital teams or retrieval
SEPSIS KILLS PROGRAM
MULTIPLE CHOICE QUESTIONS AND ANSWERS

Q1. What is sepsis?

A) Any infection
B) An infection plus hypotension
C) An infection plus systemic inflammatory response syndrome
D) Someone who is on antibiotics

Answer – C
Sepsis occurs when an infection, beginning in any tissue in the body, results in the systemic inflammatory response syndrome (SIRS). The SIRS criteria are heart rate > 90 beats/min, respiratory rate (RR) > 20 breaths/min (or partial pressure of arterial CO2 <32), temperature either > 38°C or < 36°C, and white blood cell count (WBC) either > 12,000 or < 4,000 cells/mm3. For a patient to be diagnosed with sepsis, at least two of the SIRS criteria need to be present with an infection.

Severe sepsis is defined as sepsis resulting in organ dysfunction.

Septic shock occurs when there is sepsis-induced hypotension (where either the systolic blood pressure is < 90 mmHg, 40mmHg below baseline, or the mean arterial pressure (MAP) is < 70mmHg) that persists despite adequate fluid resuscitation.


Q2. Which signs and symptoms are most common in adult patients with severe sepsis?

A) Fever > 38deg C
B) Tachycardia
C) Tachypnoea
D) Metabolic acidosis
E) Acute oliguria

Answer – C
An increase in respiratory rate is an important early marker of sepsis. Ensure that the clinical observations chart is complete and accurate.

In a large prospective cohort study of adult patients with severe sepsis (n=742), 99% had tachypnoea as defined by a respiratory rate >20. Tachycardia occurred in 97%, followed by fever (70%), acute oliguria (54%) and metabolic acidosis (38%).
(Brun-Buisson C et al. JAMA 1995)
Q3. Which of the following criteria present and new in your patient should prompt commencement of the Sepsis Pathway?

A) RR - 20 breaths/min
B) RR - 20 breaths/min + HR > 122bpm
C) RR > 25 breaths/min + HR > 120bpm + recent surgery

Answer - C
In NSW any one risk factor, sign or symptom plus 2 Yellow Zone observations or 1 Red Zone observation should prompt obtaining senior clinician review or a Rapid Response call and the consideration of the commencement of the Sepsis Pathway. Sepsis is a medical emergency.

Q4. Patients with severe sepsis and a lactate level ≥ 4 mmol/L have an increased likelihood of mortality compared to patients with a normal lactate.

A) True
B) False

Answer - True
Lactate is a normal product of anaerobic cell metabolism and is released into the blood and metabolised by the liver. Normal lactate levels are less than 1.0 mmol/L in both arterial and venous blood. Elevated serum lactate is strongly associated with morbidity and mortality in critically ill patients. One study showed a level above 4.0 mmol/L was associated with a 27% mortality rate compared with 7% for patients with a lactate of 2.5-4.0mmol/L  

(Boschert S. Is it Septic Shock? Check Lactate Level ACEP News 2007)

Elevated lactate is typically present in patients with severe sepsis or septic shock and can occur even before patients have a marked acidosis or low blood pressure. Cryptic shock is defined as a serum lactate greater than 4 mmol/L and normotensive. It is therefore important to screen for serum lactate in all patients who are suspected of sepsis irrespective of a normal blood pressure.

Elevated lactate is a late sign of sepsis in children. Where lactate is greater than 2mmol/L and sepsis is suspected, it is recommended that immediate treatment is started.

Q5. In patients with sepsis, which of the following statements is incorrect?

A) Antibiotics should never be delayed while waiting for the results of investigations.
B) Patients with febrile neutropenia should be managed using the local febrile neutropenia guideline
C) Time to first dose antibiotics does not influence outcome
D) The patient should be reviewed by the attending medical officer within 24 hours of starting on sepsis pathway and antibiotic therapy, with referral to the infectious diseases / clinical microbiology service for specific advice.

Answer - C
Intravenous antibiotic therapy should be started immediately after obtaining appropriate cultures, since early initiation of antibiotic therapy is associated with lower mortality. The choice of antibiotics should also be guided by the patient’s history including recent antibiotic exposure, comorbidities, clinical context (eg, community- or hospital-acquired infection), Gram-stain data and previous resistant bacterial isolates.
Poor outcomes are associated with inadequate or inappropriate antibiotic selection as well as delays in starting antibiotic therapy. A retrospective cohort study of 2,154 patients with septic shock found that each hour of delay in antibiotic administration over the first 6 hours was associated with an average decrease in survival of 7.6%. Therefore statement C is INCORRECT, time to first dose antibiotics DOES significantly influence outcome.

(Kumar A, Crit Care Med. 2006)

Q6. Which signs and symptoms are most common in children and infants with sepsis?

A) Fever >38deg C  
B) Persistent Tachycardia  
C) Lethargy  
D) Metabolic acidosis  
E) Lactate >4mmol/L

Answer - B

Persistent tachycardia is a reliable sign in the early recognition of sepsis in children and infants. More specifically persistent tachycardia that is resistant to treatment such as fluid bolus or antipyretics.

Not all children with sepsis will have a fever and administering antipyretics pre hospital can mask a high temperature particularly at triage and is therefore not a reliable sign or symptom of sepsis. Children can often compensate very well and relying solely on a patient’s pH again is not a reliable indicator of sepsis in children. Children and infants do not always respond to sepsis in the same way that adults do regarding an elevated lactate. It is therefore important to have a heightened level of concern with a lactate over 2mmol/L with this group.

Q7. During the monitor and reassess phase of resuscitation of the septic child or infant, which of the following signs of deterioration indicates your patient requires escalation in level of care?

A) Persistent tachycardia, slow capillary refill and hypotension  
B) Drowsiness or abnormal LOC and/or urine output<1ml/kg/hr  
C) Acidosis, increasing serum lactate or procalcitonin  
D) Hypoglycaemia, leukopenia or abnormal coagulation  
E) Any of the above

Answer - E

These are all signs suggesting your patient has severe sepsis or septic shock and has not adequately responded to treatment. This patient will require escalation in level of care and may need transfer to a paediatric Intensive Care Unit.

Q8. BP is a reliable parameter in the early recognition of sepsis in paediatric patients

A) True  
B) False

Answer- False

Children are able to maintain their BP despite significant compromise. They can prevent a reduction in their BP by vasoconstriction and increasing their heart rate. Neonates unlike adults cannot change their stroke volume so they increase their heart rate to improve cardiac output. This persistent tachycardia is a valuable sign in identifying sepsis.
Q9. Fluid resuscitation in paediatric sepsis should be given as:

A) 10ml/kg 0.9% sodium chloride over 20 minutes
B) 20ml/kg bolus of 0.9% sodium chloride over no more than 10 minutes
C) 20ml/kg bolus of 0.9% sodium chloride over 60 minutes
D) 10ml/kg 0.9% sodium chloride over 10 minutes

Answer - B

Fluid resuscitation should be given as a push of 20ml/kg 0.9% sodium chloride over no more than 10 minutes.

Q10. Key messages are: SEPSIS KILLS & TIME IS LIFE. Fill in the boxes:

- risk factors, signs, symptoms of sepsis and have early involvement of senior clinicians
- with rapid IV fluids (20ml/kg 0.9% sodium chloride) and administer antibiotics within one hour
- to appropriate in hospital teams or retrieval

Answer

RECOGNISE
RESUSCITATE,
REFER
EVALUATION

Audit tools

Sepsis database and data entry instructions
**SEPSIS DATA COLLECTION TOOL**
**ADULT INPATIENT**

<table>
<thead>
<tr>
<th>FACILITY</th>
<th>................................................</th>
</tr>
</thead>
<tbody>
<tr>
<td>WARD</td>
<td>................................................</td>
</tr>
</tbody>
</table>

For analysis purposes each facility will be able to sort for their own wards.

**Service Type**
- [ ] Critical Care
- [ ] Medical
- [ ] Surgical
- [ ] Oncology / Haematology
- [ ] Paediatric
- [ ] Aged Care
- [ ] Rehabilitation
- [ ] Maternity
- [ ] Other

**Time of recognition of sepsis**
- Rapid Response
- Clinical Review
- Other
- Time: [ ]: [ ] Date: [ ]/[ ]/[ ]

**Systolic blood pressure**
- [ ] mmHg

**Oxygen administered**
- [ ] YES
- [ ] NO
- [ ] Not required

**Lactate**
- [ ] YES
- [ ] NO
- Level: [ ] mmol/L

**Blood cultures**
- [ ] One or more sets collected
- [ ] YES
- [ ] NO

**IV resuscitation fluids**
- Administered within 60 minutes of CERS call / sepsis recognition. Only record for patients with SBP < 90mmHg and/or lactate > 4mmol/L
- [ ] Not given
- [ ] 250-500 mL
- [ ] 500-1000 mL
- [ ] > 1000 mL

**IV antibiotics**
- Administered within 60 minutes of CERS / sepsis recognition
- Do not delay for investigations or results
- First IV antibiotic commenced for sepsis:
- Time: [ ]: [ ] Date: [ ]/[ ]/[ ]

**Transfer of care**
- Use Ward if patient remaining in original ward or transferred to another ward
- Ward
- HDU / ICU
- Other hospital
- Tertiary referral hospital

**Comments**

- Signature: ........................................
- Designation: ........................................

*Mandatory Field – MRN, DOB, Lactate, IV resuscitation fluids, IV antibiotics

**THIS IS NOT A MEDICAL RECORD FORM**
COMPLETING THE DATA COLLECTION TOOL
ADULT INPATIENT

1. All dates are in dd/mm/yyyy format
2. All times are in hh:mm 24 hour clock format
3. The MRN is a MANDATORY field and cannot be modified once entered into the database.
4. The facility code originates from NSW Health and is automatically generated with your login
5. The ward code is a free text field and is for the name of the ward. Therefore for analysis of data, each facility will be able to sort on the field and monitor the progress of their own ward.
6. The service type field is used to describe the type of ward eg surgical or medical etc
7. The time and date of time of recognition of sepsis (this is the CERS call time or other method of recognition) is recorded to most accurately reflect how long it takes from time of recognition to the time of administration of the first antibiotic.
8. The Rapid Response / Clinical Review and Other boxes are to be ticked depending the call.
9. The systolic blood pressure field is the reading taken at the time of the CERS call being made / time of sepsis recognition
10. Oxygen administered is recorded as a ‘yes’ or ‘no’ if oxygen is ordered or ‘not required’ if the patient does not need oxygen.
11. Lactate is a MANDATORY field. Lactate is recorded as a ‘yes’ or ‘no’ depending on whether a level is collected. If ‘yes’ the level result is then recorded.
12. Intravenous resuscitation fluids administered if the patient has a systolic blood pressure of < 90mmHg and/or lactate > 4mmol/L at time of CERS call / sepsis recognition. Record as not given or the amount of bolus fluid given within 60 minutes of the CERS call.
13. The first intravenous (IV) antibiotic time and date is a MANDATORY field. Do not collect or enter data for patients who do not have intravenous antibiotics.
14. The transfer of care field is for where the patient was managed or was transferred to as a result of the CERS Call
15. Please use the comments box to record information which may assist in the data analysis. This includes explanations of blank fields, delays in treatment etc. Do not use the comments box as a medication prescribing or pathology record, or a medical record.
16. The signature and designation boxes are to record the name of the person completing the form.
**SEPSIS DATA COLLECTION TOOL**

**PAEDIATRIC INPATIENT**

<table>
<thead>
<tr>
<th>FACILITY</th>
<th>.........................................................</th>
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</thead>
<tbody>
<tr>
<td>WARD</td>
<td>.........................................................</td>
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</tbody>
</table>

For analysis purposes each facility will be able to sort for their own wards

<table>
<thead>
<tr>
<th>Time of recognition of sepsis</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>CERS call time / other recognition eg ward round, handover</td>
<td></td>
</tr>
</tbody>
</table>

- Rapid Response [ ]
- Clinical Review [ ]
- Other [ ]

- Time: [ ] [ ] [ ]
- Date: [ ] [ ] [ ]

<table>
<thead>
<tr>
<th>Lactate*</th>
<th>YES</th>
<th>NO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level: [ ] [ ] mmol/L</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>IV resuscitation fluid bolus*</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Administered within 60 minutes of CERS call</td>
<td></td>
</tr>
</tbody>
</table>

- Not given [ ]
- 10 mL/kg [ ]
- 20 mL/kg [ ]

<table>
<thead>
<tr>
<th>IV antibiotics *</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Administered within 60 minutes</td>
<td></td>
</tr>
<tr>
<td>Do not delay for investigations or results</td>
<td></td>
</tr>
</tbody>
</table>

- First IV antibiotic commenced for sepsis:
- Time: [ ] [ ] [ ]
- Date: [ ] [ ] [ ]

<table>
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<th>Transfer of care</th>
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<td>Use Ward if patient remaining in original ward or transferred to another ward</td>
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</tr>
</tbody>
</table>

- Ward [ ]
- ICU/HDU [ ]
- Other hospital [ ]
- Tertiary referral hospital [ ]

<table>
<thead>
<tr>
<th>Comments</th>
<th></th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Signature</th>
<th>.........................................................</th>
</tr>
</thead>
<tbody>
<tr>
<td>Designation</td>
<td>.........................................................</td>
</tr>
</tbody>
</table>

*Mandatory Field – MRN, DOB, Lactate, IV resuscitation fluids, IV antibiotics

THIS IS NOT A MEDICAL RECORD FORM
COMPLETING THE DATA COLLECTION TOOL
PAEDIATRIC INPATIENT

1. All dates are in dd/mm/yyyy format
2. All times are in hh:mm 24 hour clock format
3. The MRN is a MANDATORY field and cannot be modified once entered on the database
4. The facility code originates from NSW Health and is automatically generated with your login
5. The ward code is a free text field and is for the name of the ward. Therefore for analysis of data, each facility will be able to sort on the field and monitor the progress of their own ward
6. The time and date of time of recognition of sepsis (this is the CERS call time or other method of recognition) is recorded to most accurately reflect how long it takes from time of recognition to the time of administration of the first antibiotic
7. The Rapid Response / Clinical Review and Other boxes are to be ticked depending the call
8. Lactate is a MANDATORY field. Lactate is recorded as a ‘yes’ or ‘no’ depending on whether a level is collected. If ‘yes’ the level result is then recorded
9. Intravenous resuscitation fluid bolus administered within 60 minutes of the CERS call. Record as not given or the amount of fluid bolus given in mL/kg
10. The first intravenous (IV) antibiotic time and date is a MANDATORY field. Do not collect or enter data for patients who do not have intravenous antibiotics
11. The transfer of care field is for where the patient was managed or was transferred to as a result of the CERS Call
12. Please use the comments box to record information which may assist in the data analysis. This includes explanations of blank fields, delays in treatment etc. Do not use the comments box as a medication prescribing or pathology record, or a medical record
13. The signature and designation boxes are to record the name of the person completing the form.
<table>
<thead>
<tr>
<th>Step</th>
<th>Field Name</th>
<th>Issue</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>MRN</td>
<td>Cannot delete or change MRN once record has been generated</td>
<td>Delete record, then re-enter</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Using the tab key to move from MRN to DOB wipes out the rest of the</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>data entered</td>
<td>The MRN field has additional conditions attached to avoid entering duplicate records. You must mouse click - NOT Tab - into the next field from the MRN to allow the duplicate function to be performed</td>
</tr>
<tr>
<td>2.</td>
<td>Date and time</td>
<td>Errors in entering date and time</td>
<td>Ensure you enter the date and time in the correct order</td>
</tr>
<tr>
<td></td>
<td>• Date of birth</td>
<td></td>
<td>There are two ways to enter this data:</td>
</tr>
<tr>
<td></td>
<td>• Time of recognition</td>
<td></td>
<td>1. Mouse click into the top area and type the numbers in</td>
</tr>
<tr>
<td></td>
<td>• IV Antibiotic</td>
<td></td>
<td>2. Use the calendar. Note: you must enter the date before the time or the date of entry will default into the field</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Always check the automatically generated time calculation to ensure accuracy of entry</td>
</tr>
<tr>
<td>3.</td>
<td>Ward</td>
<td>No specific list</td>
<td>Use the name the ward is usually called</td>
</tr>
<tr>
<td>4.</td>
<td>Service Type</td>
<td>Specific services are not identified in this database</td>
<td>Use the service type that best describes your service or your patient group eg medical, surgical etc</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><em>Note: all users within your facility can see all data entered by the facility.</em></td>
</tr>
<tr>
<td>5.</td>
<td>Time of recognition of sepsis</td>
<td>Variance in time recording</td>
<td>This is the time that either a CERS call is made to escalate care of the patient, or another method of recognition i.e. patient reviewed on a ward round and care escalated after sepsis suspected</td>
</tr>
<tr>
<td>6.</td>
<td>SBP (Adults only)</td>
<td>Significant indicator of sepsis in adults</td>
<td>Particularly important in adult patients</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>This is a MANDATORY FIELD</td>
</tr>
<tr>
<td>7.</td>
<td>Oxygen administered (Adults only)</td>
<td>A component of the sepsis care bundle</td>
<td>Not all patients will require administration of oxygen</td>
</tr>
<tr>
<td></td>
<td><strong>Lactate</strong></td>
<td><strong>Significant indicator of sepsis</strong></td>
<td><strong>This is a MANDATORY FIELD</strong></td>
</tr>
<tr>
<td>---</td>
<td>-------------</td>
<td>-----------------------------------</td>
<td>-------------------------------</td>
</tr>
<tr>
<td>9.</td>
<td><strong>Blood cultures (Adults only)</strong></td>
<td>A component of the sepsis care bundle</td>
<td>Record Yes or No depending whether blood cultures are collected.</td>
</tr>
<tr>
<td>10.</td>
<td><strong>Administration of IV resuscitation fluid</strong></td>
<td>Patient meets the sepsis criteria for CERS call but IV resuscitation fluids are not prescribed / administered</td>
<td>Record as ‘not given’ if resuscitation fluid bolus not administered.</td>
</tr>
</tbody>
</table>
| 11. | **Administration of IV antibiotics** | Patient meets the sepsis criteria for CERS call but the IV antibiotics are not prescribed | If patient has not been given IV antibiotics do not record in database.  
Note: if the patient has no intravenous (IV) access, and intramuscular (IM) or intraosseous (IO) antibiotics are given instead:  
• Record time and date in this field  
• Record route of administration and reason in the comments section |
| 12. | **Transfer of care** | | Ward indicates the patient did not move treatment location as a result of sepsis recognition eg patient on ward stays on ward after sepsis recognition or patient in critical care that remains in critical care after sepsis recognition. |
| 13. | **Comments** | **Best use of this field** | **This field is not meant to be a medical record**  
Enter concise information that adds value to your data including:  
• explanation of missing data  
• hospital transfers to avoid double entries by two facilities  
• more information as thought necessary  
• specific information to identify a subset of your data for further analysis  
Use commas and spaces to separate data.  
Do not use enter key – difficult to read entry in excel spreadsheet |
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