Canadian Association of Emergency Physicians Sepsis Guidelines: the optimal management of severe sepsis in Canadian emergency departments

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CAEP Board approved June 11, 2008

CJEM 2008;10(5):443-59

ABSTRACT

Introduction: Optimal management of severe sepsis in the ED has evolved rapidly. The purpose of these guidelines is to review key management principles for Canadian emergency physicians, utilizing an evidence-based grading system.

Methods: Key areas in the management of septic patients were determined by members of the CAEP Critical Care Interest Group (C4). Members of C4 were assigned a question to be answered after literature review, based on the Oxford grading system. After completion, each section underwent a secondary review by another member of C4. A tertiary review was conducted by additional external experts, and modifications were determined by consensus. Grading was based on peer-reviewed publications only, and where evidence was insufficient to address an important topic, a “practice point” was provided based on group opinion.

Results: The project was initiated in 2005 and completed in December 2007. Key areas which were reviewed include the definition of sepsis, the use of invasive procedures, fluid resuscitation, vasopressor/inotrope use, the importance of culture acquisition in the ED, antimicrobial therapy and source control. Other areas reviewed included the use of corticosteroids, activated protein C, transfusions and mechanical ventilation.

Conclusion: Early sepsis management in the ED is paramount for optimal patient outcomes. The CAEP Critical Care Interest Group Sepsis Position Statement provides a framework to improve the ED care of this patient population.

Keywords: sepsis, severe sepsis, septic shock, emergency medicine, guidelines, resuscitation

Introduction

The management of sepsis has recently attracted long-overdue attention. Evidence supports the need for rapid identification and initiation of treatment in patients with severe sepsis. Given the widespread incidence, cost and mortality of sepsis, and because it affects patients with numerous medical and surgical comorbidities, optimal sepsis manage-
Grading System. Individual sections were combined by the (Table 1, Table 2), using the Oxford Level of Evidence members were instructed to provide objective recommendations sure all relevant studies were considered. In addition, mem-
bered to perform additional literature searches in order to en-
prise in EM, critical care medicine or both. The C4 chair de-
determined “relevant questions” to be “areas of importance for
management of severe sepsis in Canadian EM practice.” An initial PubMed search was completed by the chair using
the terms “sepsis,” “severe sepsis,” “septic shock,” “diagno-
sis,” “treatment” and “therapy,” and a reference list was
made available to C4 members using a secure Web-based portal (hosted by the Department of Emergency Medicine, Dalhousie University, Halifax, NS). Members were encour-
aged to perform additional literature searches in order to en-
sure all relevant studies were considered. In addition, mem-
ers were instructed to provide objective recommendations
to each question based on the quality of available evidence (Table 1, Table 2), using the Oxford Level of Evidence Grading System. Individual sections were combined by the chair, and a secondary review of each section was completed by C4 members. A tertiary review was completed by the en-
tire group, and all recommendations were finalized by group consensus. An additional review was also completed by ex-
ternal reviewers with expertise in sepsis. The guideline is
presented in sections: “Questions,” “Practice points,” “Rec-
ommendations” and “Rationale.” The authors have at-
tempted to base opinions on current, peer-reviewed publica-
tions. No funding from any source other than CAEP for
conference calls was used for the creation of this document. Areas that objective data do not sufficiently address, yet
were considered essential for optimal care in Canadian EM practice by C4, are presented as “Practice points.”

The authors recognize that the management of sepsis is challenging and have strived to outline complex concepts in a straightforward manner that is relevant to all Canadian EM physicians, nurses and other personnel who provide care in the ED. However, the management of sepsis will continue to evolve. As such, we anticipate that our current guidelines will require modification in the future. In addition, in order to achieve improved patient outcomes, care providers require more than knowledge of evidence-based treatments. Other key determinants include practitioner

A. Sepsis and emergency medicine

Questions

1. What is the morbidity and mortality associated with se-
vere sepsis?
2. Why is EM important in diagnosing and managing pa-
ients with sepsis?

Practice point

Severe sepsis and septic shock are common in the ED and have an associated mortality of 20%–50%. EM physicians are in a unique position to recognize and institute treat-
ments that optimize patient outcomes.

Rationale

Sepsis and septic shock are common presentations to the ED and challenge medical staff on a daily basis. In the United States, 660,000 to 750,000 cases of sepsis are estimated to occur per year with a mortality rate of approximately 20%–50%. The incidence of sepsis has increased by 8.7% per year in the last 20 years and, despite evidence that mortality rates are improving, the total number of deaths per year are increasing. In addition, patients who survive report a substantial reduction in their quality of life.

Emergency physicians are required to diagnose and manage a broad range of complaints, from those that are relatively minor in nature to those that are life-threatening. Unfortunately, the incidence of critically ill patients presenting to the ED is increasing. In addition, these patients are managed in the ED for prolonged periods of time, requiring emergency physicians to possess the knowledge and skills to effectively provide ongoing care.

Various disease states, including sepsis, have decreased morbidity and mortality as a result of early diagnosis and management. The initial hours of care for trauma patients, septic patients, myocardial infarction patients and stroke patients are critical to their long-term outcomes.

Severe sepsis and septic shock require comprehensive, aggressive and time-dependent resuscitation in the ED. Recent literature has demonstrated multiple novel options for
the management of severe sepsis and septic shock. In one study, early and aggressive resuscitation of severely septic patients in the ED resulted in a substantial improve-

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ment on mortality. Others have demonstrated increased mortality after inappropriate antimicrobial therapy selection and delay in administration. As such, the clinical management of severe sepsis and septic shock has evolved dramatically. Specific therapeutic options should be instituted in the ED to provide a morbidity and mortality benefit.

B. Definition of sepsis

Questions
1. How is sepsis defined?
2. What is the difference between sepsis, severe sepsis and septic shock?
3. What are the clinical manifestations of severe sepsis and septic shock?

Practice point
Sepsis is defined as the presence of both systemic inflammatory response syndrome and the suspicion of an infection. Sepsis is a syndrome, and can range from relatively mild (simple infection) to severe (septic shock and multi-organ dysfunction). Morbidity and mortality increase if a patient deteriorates from sepsis to severe sepsis to multi-organ dysfunction.

Rationale
Sepsis is a complex syndrome in which the clinical presentation can overlap with many other disease states. Simple bedside tests, such as serum troponins in acute myocardial infarction or bedside ultrasounds for intra-abdominal fluid, are not available. Clinicians rely on the patient history, physical exam and nonspecific laboratory investigations to diagnosis sepsis.

Sepsis has been defined as the presence, or suspicion of, an infection, in addition to evidence of increased levels of circulating inflammatory mediators, thus resulting in the

<table>
<thead>
<tr>
<th>Level</th>
<th>Therapy/prevention, etiology/harm</th>
<th>Prognosis</th>
<th>Diagnosis</th>
<th>Differential diagnosis/symptom prevalence analyses</th>
<th>Economic and decision analyses</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>SR (with homogeneity*) of RCTs</td>
<td>SR (with homogeneity*) of Level 1 diagnostic studies; CDR† with 1b studies from different clinical centres</td>
<td>SR (with homogeneity*) of prospective cohort studies</td>
<td>SR (with homogeneity*) of Level 1 economic studies</td>
<td></td>
</tr>
<tr>
<td>1b</td>
<td>Individual RCT (with narrow confidence interval†)</td>
<td>Individual inception cohort study with ≥ 80% follow-up; CDR† validated in a single population</td>
<td>Validation of Level &gt; 2 cohort study with good reference standards or CDR† tested within one clinical centre</td>
<td>Prospective cohort study with good follow-up for Level &gt; 2 cohort studies</td>
<td>Analysis based on clinically sensible costs or alternatives; systematic review(s) of the evidence; and including multi-way sensitivity analyses</td>
</tr>
<tr>
<td>1c</td>
<td>All or none§</td>
<td>All or none case-series</td>
<td>Absolute SpPins and SnNouts</td>
<td>All or none case-series</td>
<td>Absolute better-value or worse-value analyses for Level &gt; 2 cohort studies</td>
</tr>
<tr>
<td>2a</td>
<td>SR (with homogeneity*) of cohort studies</td>
<td>SR (with homogeneity*) of Level &gt; 2 diagnostic studies</td>
<td>SR (with homogeneity*) of 2b and better studies</td>
<td>SR (with homogeneity*) of Level &gt; 2 economic studies</td>
<td></td>
</tr>
<tr>
<td>2b</td>
<td>Individual cohort study (including low quality RCT; e.g., &lt; 80% follow-up)</td>
<td>Retrospective cohort study or follow-up of untreated control patients in an RCT; derivation of CDR† or validated on split-sample only</td>
<td>Retrospective cohort study, or poor follow-up</td>
<td>Analysis based on clinically sensible costs or alternatives; limited review(s) of the evidence, or single studies; and including multi-way sensitivity analyses</td>
<td></td>
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<tr>
<td>2c</td>
<td><em>Outcomes</em> research; ecological studies</td>
<td><em>Outcomes</em> research</td>
<td>Ecological studies</td>
<td>Audits or outcomes research</td>
<td></td>
</tr>
<tr>
<td>3a</td>
<td>SR (with homogeneity*) of case-control studies</td>
<td>SR (with homogeneity*) of 3b and better studies</td>
<td>SR (with homogeneity*) of 3b and better studies</td>
<td>SR (with homogeneity*) of 3b and better studies</td>
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</tr>
</tbody>
</table>

Continued on next page
systemic inflammatory response syndrome.22 Circulating inflammatory mediators may cause a deviation of patients’ core body temperature, heart rate, respiratory rate and white blood cell count.23,24 However, inflammatory mediators can be caused by pathological states other than sepsis, such as thermal injury, pancreatitis, acute thyroiditis and traumatic injury.23 The clinical syndrome characterized by a noninfective systemic inflammatory response syndrome reaction is similar to those of a septic patient.23,24 The only difference is that, in sepsis, micro-organisms have invaded a sterile body tissue. Others have recently revisited the definition of sepsis with a goal to produce a definition that is usable both for the bedside clinician and for research.22

Clinical recognition requires physician knowledge and skill, as septic patients present to the ED with a wide spectrum of illness. Traditionally, 2 or more of the following variables have been used (coupled with evidence/presumption of an infection) to identify patients who are septic,22,25

1. Body temperature less than 36°C or greater than 38°C.
2. A heart rate greater than 90 beats per minute.
3. A respiratory rate greater than 20 breaths per minute, PACO₂ less than 30 mm Hg or the need for mechanical ventilation.
4. A white count less than 4000 or greater than 12 000, or greater than 10% immature bands.

Severe sepsis is defined as sepsis in addition to evidence of end organ dysfunction. Virtually any organ system may be affected in severe sepsis.22 Organ system dysfunctions

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**Table 1. Oxford Centre for Evidence-based Medicine Levels of Evidence (May 2001) (Part 2 of 2)**

<table>
<thead>
<tr>
<th>Level</th>
<th>Therapy/prevention, etiology/harm</th>
<th>Prognosis</th>
<th>Diagnosis</th>
<th>Differential diagnosis/symptom prevalence study</th>
<th>Economic and decision analyses</th>
</tr>
</thead>
<tbody>
<tr>
<td>3b</td>
<td>Individual case-control study</td>
<td></td>
<td>Non-consecutive study; or without consistently applied reference standards</td>
<td>Non-consecutive cohort study, or very limited population</td>
<td>Analysis based on limited alternatives or costs, poor quality estimates of data, but including sensitivity analyses incorporating clinically sensible variations.</td>
</tr>
<tr>
<td>4</td>
<td>Case-series (and poor quality cohort and case-control studies§§)</td>
<td>Case-series (and poor quality prognostic cohort studies****)</td>
<td>Case-control study, poor or non-independent reference standard</td>
<td>Case-series or superseded reference standards</td>
<td>Analysis with no sensitivity analysis</td>
</tr>
<tr>
<td>5</td>
<td>Expert opinion without explicit critical appraisal, or based on physiology, bench research or “first principles”</td>
<td>Expert opinion without explicit critical appraisal, or based on physiology, bench research or “first principles”</td>
<td>Expert opinion without explicit critical appraisal, or based on physiology, bench research or “first principles”</td>
<td>Expert opinion without explicit critical appraisal, or based on physiology, bench research or “first principles”</td>
<td></td>
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</tbody>
</table>

Notes: Users can add a minus-sign “−” to denote the level of that fails to provide a conclusive answer because of
1. EITHER a single result with a wide confidence interval (such that, for example, an ARR in an RCT is not statistically significant but whose confidence intervals fail to exclude clinically important benefit or harm);
2. OR a systematic review with troublesome (and statistically significant) heterogeneity;
3. Such evidence is inconclusive, and therefore can only generate Grade D recommendations.

* By homogeneity we mean a systematic review that is free of worrisome variations (heterogeneity) in the directions and degrees of results between individual studies. Not all systematic reviews with statistically significant heterogeneity need be worrisome, and not all worrisome heterogeneity need be statistically significant. As noted above, studies displaying worrisome heterogeneity should be tagged with a “−” at the end of their designated level.

† Clinical decision rule. (These are algorithms or scoring systems which lead to a prognostic estimation or a diagnostic category.)
†† See note #2 for advice on how to understand, rate and use trials or other studies with wide confidence intervals.
§ Met when all patients died before the Rx became available, but some now survive on it; or when some patients died before the Rx became available, but none now die on it.
§§ By poor quality cohort study we mean one that failed to clearly define comparison groups and/or failed to measure exposures and outcomes in the same (preferably blinded), objective way in both exposed and non-exposed individuals and/or failed to identify or appropriately control known confounders and/or failed to carry out a sufficiently long and complete follow-up of patients. By poor quality case-control study we mean one that failed to clearly define comparison groups and/or failed to measure exposures and outcomes in the same (preferably blinded), objective way in both cases and controls and/or failed to identify or appropriately control known confounders.
*** Split sample validation is achieved by collecting all the information in a single tranche, then artificially dividing this into “derivation” and “validation” samples.
**** An “Absolute SpPin” is a diagnostic finding whose Specificity is so high that a Positive result rules in the diagnosis. An “Absolute SnNOut” is a diagnostic finding whose Sensitivity is so high that a Negative result rules out the diagnosis.
††† Good, better, bad and worse refer to the comparisons between treatments in terms of their clinical risks and benefits.
††‡‡ Good reference standards are independent of the test, and applied blindly or objectively to applied to all patients. Poor reference standards are haphazardly applied, but still independent of the test. Use of a non-independent reference standard (where the ‘test’ is included in the ‘reference’, or where the ‘testing’ affects the ‘reference’) implies a level 4 study.
†††† Better-value treatments are clearly as good but cheaper, or better at the same or reduced cost. Worse-value treatments are as good and more expensive, or worse and the equally or more expensive.

** Validating studies test the quality of a specific diagnostic test, based on prior evidence. An exploratory study collects information and trawls the data (e.g. using a regression analysis) to find which factors are ‘significant’.
** By poor quality prognostic cohort study we mean one in which sampling was biased in favour of patients who already had the target outcome, or the measurement of outcomes was accomplished in < 80% of study patients, or outcomes were determined in an unblinded, non-objective way, or there was no control for confounding factors.

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include the central nervous system (agitation, depressed level of consciousness), the cardiovascular system (tachycardia, hypotension, systemic vasodilatation), the gastrointestinal system (elevated liver enzymes), the renal system (elevated creatinine, low urine output) and the hematological system (coagulation dysfunction, thrombocytopenia). Once a patient progresses from sepsis to severe sepsis or septic shock, his or her risk of mortality increases substantially.

Septic shock is differentiated from severe sepsis by the presence of ongoing cardiovascular instability despite adequate fluid resuscitation. Systolic blood pressures under 90 mm Hg are considered insufficient for optimal cellular homeostasis in septic patients. It is common for septic shock patients to require vasopressor medication infusions for stabilization of the systemic vasodilatation resulting from widespread cytokine release.

The definition of sepsis will continue to evolve as new diagnostic tests are identified to aid in diagnosis. In fact, it is now recognized that patients can be in septic shock without hemodynamic instability (“cryptic shock”). At the present time, emergency physicians should be familiar with the current definition in order to implement beneficial management options.

C. The pathophysiology of sepsis

Question

What should emergency physicians understand about the pathophysiology of sepsis?

Practice point

A patient’s immune response to microbes results in a myriad of effects that are not limited to the site of infection. Neurohumoral response and vascular cytokine release may result in various organ dysfunctions.

Rationale

Sepsis is a complex physiological process. Microscopically, once infection is introduced to a host, an initial neurohumoral response and release of cytokines activates inflammatory mediators. Additional responses include neutrophil adhesion to the endothelium, endothelial disruption and activation of clotting. When combined, these effects contribute to procoagulation and microthromboses at the cellular level. Excess inflammation and microthromboses lead to cellular hypoxia, anaerobic metabolism and cell death.

On a macroscopic level, sepsis progresses along a spectrum from infectious insult to septic shock and multiorgan dysfunction. Septic shock is a form of distributive shock, manifested in patients by an increase in cardiac output and vasodilatation. The oxygen demand of end organs exceeds oxygen delivery, and with the cellular switch from aerobic to anaerobic metabolism, lactate is produced. As oxygen demands remain unmatched, perfusion decreases leading to end organ failure. With each organ system failure, absolute mortality increases by 15%–20%.

D. Invasive procedures in septic ED patients

Questions

1. When is endotracheal intubation indicated in septic patients?
2. When does a septic patient require a central venous line?
3. What location should be used when inserting a central line?
4. Which patients require an arterial line?

Recommendations

Endotracheal intubation should be instituted when required for airway protection, support of oxygenation or assisted ventilation. (Grade D)

Patients with severe sepsis, sepsis-induced tissue hypoperfusion or hypotension unresponsive to initial fluid resuscitation should have central venous access. (Grade D)

The internal jugular or subclavian veins are the preferred sites for central venous cannulation. (Grade D)

All patients who are receiving vasopressor support should have an arterial line inserted as soon as resources allow. Arterial lines should be strongly considered in patients who are hemodynamically labile or who require frequent blood work and do not have central venous access. (Grade D)

Rationale

There is little evidence that most invasive procedures influence the outcome of septic patients. Recommendations are based on basic principles and common practice in emergency medicine and critical care medicine.

Table 2. Grades of Recommendation

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
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<tbody>
<tr>
<td>A</td>
<td>consistent level 1 studies</td>
</tr>
<tr>
<td>B</td>
<td>consistent level 2 or 3 studies or extrapolations from level 1 studies</td>
</tr>
<tr>
<td>C</td>
<td>level 4 studies or extrapolations from level 2 or 3 studies</td>
</tr>
<tr>
<td>D</td>
<td>level 5 evidence or troublingly inconsistent or inconclusive studies of any level</td>
</tr>
</tbody>
</table>

Note: “Extrapolations” are where data is used in a situation that has potentially clinically important differences than the original study situation. Reprinted with permission from the Oxford Centre for Evidence-Based Medicine. www.cebm.net/index.aspx?o=1025
Indications for endotracheal intubation are airway protection, or the support of ventilation (CO₂ clearance) or oxygenation. These situations are common in critically ill septic patients. Positive pressure ventilation can reduce venous return, particularly in patients with an intravascular fluid deficit, and thus decrease blood pressure. Fluid resuscitation should occur prior to and in a simultaneous fashion with intubation to mitigate this effect. Conversely, positive pressure ventilation, in association with sedation, has beneficial effects on hemodynamics by reducing afterload and relieving the effect of work of breathing in the adequately volume-resuscitated patient.³⁹

Central venous access in the septic patient may be indicated for the measurement of venous oxygen saturation or central venous pressure or for the infusion of vasoactive medications. A landmark trial of ED sepsis demonstrated that patients who received early goal-directed therapy had a lower mortality rate.³² In this study, there were multiple therapeutic interventions given to the treatment group, all with the goal of rapidly improving tissue perfusion. It is unclear which component(s) of the treatment algorithm provided benefit. The “early goal-directed therapy” treatment bundle included the use of central venous oxygen saturation (necessitating the insertion of a central venous catheter) and targeted initial fluid resuscitation to a central venous pressure greater than or equal to 8–12 mm Hg.

Approximately 50% of septic patients who initially present with severe sepsis will respond to fluids alone.³⁰ It is reasonable to perform initial fluid resuscitation targeting clinical end points, such as blood pressure, heart rate and urine output. In situations where there is uncertainty about the adequacy of fluid restoration or concern of fluid overload, central venous monitoring should be started.

Central venous access is usually achieved with placement of subclavian, internal jugular or femoral vein catheters. Venous sampling from these lines is a measure of the mixed venous oxygen saturation and can be used as a surrogate marker of tissue oxygen delivery and extraction. The closer the sample site is to the right ventricle, the more closely it will represent a true mixed venous sample.³¹ Femoral venous samples from septic patients may overestimate the drop in venous saturation, but values from this site remain useful, particularly when trended to evaluate the effectiveness of therapy.

Arterial catheters provide continuous blood-pressure monitoring and facilitate blood sampling. Patients who require vasopressor or inotropic support should have an arterial line inserted as soon as the clinical situation allows, enabling medication titration and monitoring.³² Frequent cuff blood-pressure measurement is a reasonable alternative in some clinical situations.

E. Fluid resuscitation in septic ED patients

Questions

1. What are the physiologic goals of fluid resuscitation in severe sepsis?
2. What type of fluid should be used to resuscitate septic patients?
3. How much is too much? Can I overresuscitate a patient with septic shock?

Recommendation

Fluid resuscitation in septic ED patients with tissue hypoperfusion should be initiated immediately upon recognition. (Grade B)

There is currently no conclusive evidence supporting the type of fluid used in resuscitation of the septic patient. Reasonable choices include Ringer’s lactate, normal saline or albumin. (Grade D)

An initial bolus of 1–2 L of crystalloid or 500–1000 mL of colloid should be given over 30–60 minutes and repeated as required to correct tissue perfusion and/or blood pressure abnormalities. (Grade D)

Initial fluid resuscitation should target clinical response, but in patients where there is concern about fluid overload, uncertainty about the adequacy of volume resuscitation or suboptimal response, fluid therapy should be guided by central venous pressure monitoring, targeting a pressure of 8–12 mm Hg. (Grade D)

Rationale

Septic patients may have a relative intravascular fluid deficit as a result of vasodilation and the loss of intravascular fluid because of an increase in microvascular permeability. In addition, these derangements are often accompanied by fluid losses due to vomiting, diarrhea, diaphoresis, fever, a reduction in fluid intake or bleeding.

The first priority in treating decreased tissue perfusion or hypotension is rapid fluid replacement. Intravascular fluid replacement increases cardiac preload, resulting in higher stroke volume, increased cardiac output and improved organ perfusion.

The volume of fluid administered will depend on patient factors, the progression of the disease state and the type of fluid infused. Rackow found that septic patients often required up to 8 L of crystalloid or 3 L of colloid in the first 24 hours.³³ Patients in the Early Goal-Directed Therapy Trial received an average of 5 L of fluid in the first
6 hours. Fluids should be administered aggressively, targeting clinical signs of improved tissue perfusion, and, if possible, a central venous pressure of 8–12 mm Hg.

The central venous pressure target of 8–12 mm Hg has not been demonstrated to be superior to higher or lower values, but is based on physiologic principles. It is sufficiently high to ensure adequate preload but generally not high enough to promote pulmonary edema formation. Patients undergoing positive pressure ventilation may require higher filling pressures.

Clinicians are often concerned that aggressive intravenous fluid administration will result in pulmonary edema and hypoxia. The majority of septic patients with hypoxia secondary to interstitial fluid accumulation are not truly volume-overloaded; rather, they are suffering from an alveolar capillary leak syndrome. Fluid resuscitation must be tailored to the clinical situation, balancing volume needs and oxygenation to maximize oxygen delivery at the tissue level.

Randomized controlled trials and systematic reviews about the choice of crystalloid or colloid in critically ill patients have not demonstrated a single ideal fluid. A large trial of 6997 critically ill patients found no mortality or morbidity difference between normal saline and 4% albumin. A recent comparison of 10% pentastarch versus Ringer’s lactate that focused on severe sepsis patients also found no mortality difference but did report an increase in acute renal failure in the 10% pentastarch group. Until further research is completed on the use of starch containing fluids, it would be prudent to avoid their use in septic patients. Overall, the choice should be made based on availability, cost and patient needs. For most patients in the ED, resuscitation with normal saline or lactated Ringer’s solutions may be the most appropriate strategy.

**F. Vasopressor and inotrope infusions in the ED**

**Question**

1. What is the difference between vasopressors and inotropes?

**Practice point**

Vasopressor medications increase vascular constriction via peripheral α receptors and therefore increase blood pressure. Inotrope medications increase myocardial contractility and heart rate via β receptor stimulation but may not increase blood pressure.

**Rationale**

Vasopressors constrict blood vessels, which, in turn, elevates blood pressure. This vasoconstriction is considered beneficial by increasing the perfusion pressure to various tissue beds. Conversely, in some patients vasopressors may increase myocardial afterload and also reduce flow in some vascular beds (e.g., splanchnic). Examples of vasopressors include phenylephrine, norepinephrine and epinephrine.

Inotropes increase cardiac contractility and cardiac output. Commonly used inotropes include dobutamine and milrinone. These drugs will not necessarily increase blood pressure and can lead to a decrease in blood pressure caused by peripheral dilation. Indeed, if the blood pressure is low, it is often necessary to administer an inotrope in combination with a vasopressor to maintain mean arterial pressure. Some medications are a combined vasopressor-inotrope, as they contain properties of both (e.g., epinephrine, norepinephrine and dopamine).

**Question**

2. When should vasopressors be used?

**Recommendations**

After adequate volume resuscitation, vasopressor medications should be used to maintain blood pressure (mean arterial blood pressure [MAP] > 65 mm Hg) and organ perfusion. (Grade B)

Vasopressor medications may be required to maintain blood pressure prior to adequate volume resuscitation. (Grade D)

**Rationale**

Many patients in septic shock who are volume resuscitated (i.e., central venous pressure [CVP] 8–12) remain hypotensive or inadequately perfused. For these patients, vasopressors are recommended to achieve a MAP of greater than 65 mm Hg. Other indicators of organ perfusion should be closely monitored, including the level of consciousness, peripheral perfusion and urine output. In most individuals, a urine output of 0.5 mL/kg/h is acceptable.

In the ED, patients in septic shock are often initially hypovolemic and hypotensive. Vasopressors may be used concurrently with volume resuscitation to maintain blood pressure until adequate volume status can be established. Once the patient is volume resuscitated, vasopressor infusion should be reduced if physiologic goals are achieved.

**Question**

3. Which vasopressor should be used?

**Recommendation**

Various vasopressors, including norepinephrine and dopamine, may be used in septic shock. (Grade C)
Rationale

Insufficient data is available to definitively conclude which optimal vasopressor should be used in septic patients. However, there is sufficient data to recommend the use of norepinephrine (levophed) and dopamine as first-line agents.45,47-49 Both norepinephrine and dopamine increase MAP in volume-replete patients in septic shock.32,44,45,47,48 In addition, there is evidence that norepinephrine may increase MAP in patients who are refractory to the administration of intravenous fluids and dopamine.45,51-57 However, dopamine has been in widespread use in the ED for decades and is conveniently supplied in a premixed solution.

Recommended dopamine doses range from 1 to 25 µg/kg/min.45 Low-dose dopamine does not protect critically ill patients from renal dysfunction and infusions at that rate for that purpose should no longer be used.58,59 Low-dose dopamine (< 5 µg/kg/min) may increase urine output but does not reduce multiorgan dysfunction or the need for renal replacement therapy.58,60 Norepinephrine should be titrated based on patient response, including blood pressure and urinary output.32,45 Norepinephrine does not worsen renal dysfunction in septic patients.49,55,57

Other vasopressors, such as epinephrine and vasopressin, may also be used in septic patients, although current practice in Canadian EDs is limited. Vasopressin is a potent vasoconstrictor that acts as an agonist at the V1 receptor to constrict vascular smooth muscle. It may be used as an adjunct to decrease the dose of other vasopressors in septic shock.61-63 Epinephrine increases mean arterial pressure, but concerns exist as it may impair splanchnic circulation and raise lactate levels in septic patients.47

Question

4. How should vasopressors be administered in the ED (central v. peripheral)?

Practice point

Vaspressors should be administered through a central venous line in the ED.

Rationale

It may be necessary to deliver vasopressors peripherally until a central line can be established in the ED. While vasopressors can be delivered peripherally in a temporary fashion, the risks of this practice include vein sclerosis, extravasation and inadequate or uncertain delivery.64-67 Central access should be established in all septic shock patients as soon as possible, given the competing clinical demands of the department. Central-line insertion also allows for rapid fluid infusion and central venous oxygen saturation monitoring.

Question

5. When should inotropes be used?

Recommendation

Inotropes should be used to maintain central venous oxygen saturation over 70% (Grade D) and adequate cardiac output (Grade D) in volume-replete septic patients who have an adequate blood pressure.

Rationale

A pivotal study of early goal-directed therapy in the ED demonstrated the utility of optimal saturation of the venous blood returning to the right ventricle after being deoxygenated in peripheral tissues.12 Inotropes should be titrated to maintain central venous oxygen saturation greater than 70% (a measure of adequate resuscitation) as part of early goal-directed therapy in septic shock.12 Specifically, these patients should have a CVP of 8–12 mm Hg, a mean arterial pressure greater than 65 mm Hg, adequate urine output and a hematocrit (Hct) over 30%, prior to initiation of inotropes to improve oxygen delivery.12,44,45 This therapy usually requires a subclavian or an internal jugular central line for appropriate monitoring and titration. Ideally, inotropes should not be titrated to blood pressure alone, but should include ongoing assessment of the patient’s overall progress.

Question

6. Which inotrope should be used?

Recommendation

 Dobutamine is considered first line in septic shock patients who require inotropic support. (Grade D)

Rationale

Inotropes have been studied less extensively than vasopressors in septic shock. Dobutamine, in doses of 2–28 µg/kg/min, effectively raises cardiac output in this population.45,68 However, caution is warranted, as tachycardia and hypotension are common in patients receiving dobutamine. The avoidance of further hypotension is important and, therefore, patients often require additional vasopressor infusions when dobutamine is required. Milrinone infusion may be considered if tachycardia limits the use of dobutamine.

G. Microbial cultures in the ED

Question

1. What is the importance of cultures in septic ED patients?
Recommendation
The establishment of a microbiological diagnosis and confirmation of effective antimicrobial therapy requires cultures of all potentially infected sites. (Grade D)

Rationale
Data on the benefit of body fluid cultures in septic ED patients is incomplete. Unfortunately, cultures are often negative, and the yield is influenced by the source of infection and type of culture. For example, one series states that no more than 50% of patients with severe sepsis will have positive blood cultures. However, as a “standard-of-care,” patients with possible severe sepsis should have all potential sources of infection cultured. Cultures maximize information to aid in the establishment of a definitive microbiological diagnosis, utilizing appropriate antibiotics and providing data about local pathogens and antibiotic resistance.

Question
2. When should cultures be obtained in septic emergency patients?

Recommendation
Cultures should be obtained as soon as a patient is identified as potentially septic. Ideally, this should be done prior to antimicrobial administration, but should not delay treatment. (Grade D)

Rationale
Insufficient data exists on the timing of culture acquisition to definitively answer this question. Regardless, it is generally accepted that cultures should be obtained prior to the initiation of antibiotic therapy. This offers the best yield, and assists in the re-evaluation of antimicrobials when cultures results are available 24–72 hours later. However, given mounting evidence that delaying antibiotic therapy is harmful (see section I. Antimicrobial administration in early sepsis), cultures should be drawn as soon as possible and should not delay treatment.

Question
3. How and where should cultures be obtained?

Recommendation
Cultures should be obtained from all potential sources of infection. (Grade D)

Rationale
Cultures drawn in septic ED patients should be dictated by the suspected source of infection. Usual practice is to obtain blood cultures from all potentially infected sites. This typically means peripheral blood, urine and sputum cultures. Other sources, such as cerebrospinal fluid, cutaneous wounds and indwelling vascular access devices should also be cultured based on clinical suspicion. The use of standard protocol for obtaining blood cultures can result in better yield and less contamination. To maximize yield, a minimum of 20 mL per blood culture is recommended (2 bottles of 10 mL each).

H. Bedside prognostication of patients with sepsis

Question
Are there bedside predictors of outcome in septic patients?

Recommendation
The elevation of serum lactate in patients with sepsis is associated with an increase in mortality. Organ dysfunction also indicates an increased risk of death. (Grade B)

Rationale
Lactate is generated during anaerobic cellular metabolism and is a hallmark of severe sepsis or septic shock. Lactate has been studied in a variety of clinical conditions in critical care medicine and has been demonstrated to predict mortality with varied accuracy. Original work in the 1960s in patients with shock and elevated lactates (> 4 mmol/L) demonstrated a mortality rate greater than 80%. Bernardin also found that a lactate of > 3.5 mmol/L at 24 hours after admission was associated with elevated mortality (approximately 30%). Another trial of 1228 patients with infection-related diagnoses found that a lactate of greater than 4 mmol/L was associated with a mortality of 22% within 3 days, versus 2% for patients with a lactate of less than 4 mmol/L. Conversely, patients in whom serum lactate decreases after ED therapy have an associated reduction in mortality. Although lactate may be generated by various mechanisms, the elevation of serum lactate in a patient with septic shock indicates that emergent resuscitation is required. Early measurement of serum lactate may aid in the identification of patients where the severity of the infection may otherwise be underappreciated.

Organ dysfunction in patients with severe sepsis also predicts mortality. An ED-based observational cohort study of 3102 patients with suspected infection demonstrated a 4-fold increase of in-hospital mortality in patients with severe sepsis (failure of 1 or more organs) versus patients without organ dysfunction. An increase in mortality risk was demonstrated as the number of dysfunctional organs increased. In patients with septic shock...
on ED presentation and organ dysfunction, the risk of death was 13 times higher than in septic patients without organ dysfunction.\textsuperscript{78}

\textit{I. Antimicrobial administration in early sepsis}

\textbf{Question}

1. When should antibiotics be administered?

\textbf{Recommendation}

Patients presenting with severe sepsis/septic shock should receive broad spectrum antibiotics as soon as possible, ideally within 60 minutes of the recognition of severe sepsis. (Grade C)

\textbf{Rationale}

Rapid administration of antibiotics to patients with severe sepsis/septic shock should be considered of paramount importance. Animal models suggest that the administration of antibiotics within 12 hours resulted in a mortality rate of 20\% versus over 85\% if delayed for more than 15 hours.\textsuperscript{79} Prospective human trials in severe sepsis/septic shock are lacking, but studies in meningitis suggest delaying antibiotics over one hour is associated with increased mortality.\textsuperscript{80} Patients with community-acquired pneumonia also had similar reductions in mortality. Houck and colleagues\textsuperscript{81} found that those patients who received antibiotics within 4 hours of arrival had lower mortality (11.6\% v. 12.7\%), and decreased length of hospital stay. Kumar and coauthors\textsuperscript{82} retrospectively studied over 2000 patients with septic shock. Patients who received antibiotics within the first hour had a mortality of 20\%, and delayed antimicrobial administration was associated with increased mortality. The study concluded that timing of antibiotic administration was the most significant determinant of outcome and that for every hour delay following onset of hypotension mortality increased by 8\%.\textsuperscript{82} In addition, only 50\% of patients in this study received antibiotics within the first 6 hours of diagnosis of septic shock in the ED. Measures to ensure that antimicrobials are administered immediately by a member of the health care team are recommended.

\textbf{Question}

2. What antibiotics should be administered?

\textbf{Recommendation}

The administration of appropriate antimicrobials should provide appropriate coverage against all likely pathogens and should be based on patient characteristics and local epidemiology and resistance patterns. (Grade B)

\textbf{Rationale}

It is well established that appropriate initial antimicrobial therapy in sepsis reduces morbidity and mortality. Multiple studies in patients with gram negative bacteremia have demonstrated an approximate 50\% reduction in mortality with appropriate antimicrobial therapy.\textsuperscript{18-31} Specific antimicrobial recommendations are beyond the scope of this document but several published recommendations exist.\textsuperscript{83,84} Emphasis should be on initial broad-spectrum coverage and should reflect the likely source and the local resistance patterns. Either monotherapy or dual therapy in uncomplicated cases is acceptable based on available data.

\textbf{Question}

3. Are there special circumstances that modify antimicrobial recommendations in ED patients?

\textbf{Recommendation}

In patients with neutropenia or who are immunocompromised, the possibility of infection with \textit{Pseudomonas} sp. must be considered. Coverage with 2 antimicrobial agents effective against this organism should be considered. (Grade D)

\textbf{Rationale}

Immunocompromised patients are at high risk of serious infection with \textit{Pseudomonas} sp. Controversy remains as to the benefit of prolonged dual therapy versus monotherapy, but it is generally agreed that initial dual therapy coverage is appropriate for serious infection.\textsuperscript{83,84} The Infectious Diseases Society of America recently published guidelines that address this issue.\textsuperscript{85}

Other special circumstances include nursing home patients, patients recently discharged from a health care facility and those recently (< 3 mo) or currently receiving antimicrobial medications. EM physicians should consider these factors when deciding on appropriate antimicrobial therapy.

\textbf{J. Source control in sepsis}

\textbf{Question}

Is source control important in patients with severe sepsis?

\textbf{Recommendation}

Optimal treatment includes consideration of all potential sources of infection and their subsequent control or elimination. As with antimicrobials, source control should be a priority in ED management. (Grade D)
**Rationale**

Source control should be a primary focus in the initial management of septic patients in the ED. Unfortunately, the identification and removal of a focus of infection may often be overlooked. Examples include removing potentially infected venous catheters, draining abscesses and debriding necrotic tissue (such as in necrotizing fasciitis). Recommendations are largely based upon expert opinion rather than randomized trials. Several case series and case reports have suggested the superiority of early source control versus late or no control at all. For example, one small case series reporting on children with necrotizing fasciitis found that delayed debridement of infected tissue was associated with high mortality rates. As such, source control should occur as soon as possible.

**K. The use of activated protein C (APC) in the ED**

**Question**
What is the role of APC in the ED?

**Recommendation**

APC infusions may be considered in adult patients with severe sepsis (Acute Physiology and Chronic Health Evaluation [APACHE] II ≥ 25, or 2 or more organ dysfunctions) within 24 hours of diagnosis. (Grade B)

Little data is available on the use of APC in the ED.

**Rationale**

APC is an anticoagulant that combats the hypercoagulable state associated with sepsis by various mechanisms. Administration requires a weight-based, 96-hour infusion protocol that is combined with other standard supportive care measures. Based on a prospective randomized controlled trial of 1690 patients with severe sepsis that demonstrated a 6.1% absolute mortality reduction, APC is used in the management of severely septic patients.

In the United States, APC received an indication from the Food and Drug Administration for the treatment of severe sepsis in patients with an APACHE II score of 25 or greater. Most Canadian health jurisdictions have adopted similar indications, although some centres instead use greater than 2 organ system dysfunction as an indication. Surgical patients and those with active bleeding or a significant risk of blood loss are not suitable candidates for APC, as the risk of hemorrhage is increased. In addition, APC is not indicated for patients with sepsis who have a low risk of death (APACHE II score < 25 or ≤ 1 organ dysfunction) or for children.

Data on the use of APC in an ED setting is incomplete. To date, there exists no placebo-controlled, double-blind, randomized controlled trial specifically examining the issue of timing of APC administration. Post hoc analysis of available data indicates the potential for improved outcome with earlier administration in select patients.

Since the use of APC is based on septic patients having an APACHE II score of 25, and with limited data on its use in the ED, APC should not be routinely administered in the ED, unless under the direction of an intensivist or infectious disease specialist.

**L. Corticosteroid use in septic ED patients**

**Question**

1. When should corticosteroids be used in septic ED patients?

**Recommendation**

Hemodynamically unstable patients who do not respond to volume resuscitation and vasopressor infusion may be considered for low-dose corticosteroid replacement in the ED. (Grade D)

**Rationale**

The use of low-dose corticosteroids in severe sepsis is supported by recent trials and meta-analyses, although significant controversy remains. Annane and co-authors randomized vasopressor-dependent septic patients to receive either placebo or hydrocortisone 50 mg intravenously (IV) every 6 hours and fludrocortisone 50 µg orally once daily. Patients were also administered a 0.25-mg adrenocorticotropic hormone (ACTH) stimulation test in both groups to determine the adrenal response to direct stimulation. Among nonresponders to ACTH, 28-day mortality was significantly reduced in the steroid group at 53% versus 63% for the control group (p = 0.04). There was no statistical difference in corticosteroid-related complications, such as additional infections or gastrointestinal bleeding.

Other authors have also investigated the utility of corticosteroids in septic patients. Meta-analyses concluded a benefit for the administration of low-dose corticosteroids (200–300 mg/d of hydrocortisone equivalents) in ICU patients with septic shock. Annane and colleagues reviewed trials involving low-dose corticosteroid for at least 5 days found a significant decrease in mortality (relative risk [RR] 0.80, 95% confidence interval [CI] 0.67–0.95, p = 0.01). In addition, a meta-analysis performed by Minneci and colleagues determined that low-dose glucocorticoids provided a beneficial effect on...
survival (RR 1.23, 95% CI 1.01–1.50, \( p = 0.036 \)) and shock reversal (RR 1.71, 95% CI 1.29–2.26, \( p < 0.001 \)).

However, the corticosteroid therapy of septic shock (CORTICUS) trial, which randomized 499 severe septic patients to receive corticosteroids or placebo, failed to confirm a benefit of steroids in sepsis.\(^9\) In this study, mortality was not affected by the use of corticosteroids despite more rapid resolution of hemodynamic instability in patients receiving corticosteroids. In addition, there was an increased incidence of infectious complications in patients who received corticosteroids. Unfortunately, this study suffered from a lack of power because of its premature termination secondary to poor recruitment and the loss of funding. In addition, etomidate use in patients requiring intubation was identified as a possible confounder, as it has been associated with adrenal insufficiency.

Based on the available evidence, corticosteroids may be considered in patients with severe sepsis and hemodynamic instability. The utility of corticosteroids remains unproven.

**Question**

2. How should steroids be used in patients with septic shock?

**Recommendation**

Doses of 200–300 mg per day of hydrocortisone may be considered in patients requiring vasopressor support. (Grade D)

**Rationale**

Low-dose corticosteroid replacement may be considered in septic shock patients. With the exception of 1 trial,\(^\) the entry criteria for studies involving low-dose corticosteroids required the use of vasopressor support.\(^16,93-95\) Annane and colleagues\(^16\) required a systolic blood pressure less than 90 mm Hg for at least 1 hour despite adequate fluid replacement and the use of dopamine at a dose greater than 5 \( \mu \)g/kg/min or current treatment with norepinephrine or epinephrine.

Hydrocortisone may be used for glucocorticoid replacement, in doses of 200–300 mg/d in divided (every 6–8 h) doses. The use of additional mineralocorticoid replacement is controversial.

The considerable heterogeneity in the use of the ACTH stimulation test\(^100-104 \) led a recent review to conclude that the ACTH test is optional.\(^105\) In addition, some centres may take several days to return ACTH stimulation results, which may reduce the relevance of this test. The administration of corticosteroids should not be delayed pending the results of the ACTH test. Rather, the ACTH test should be performed and blood sent to the receiving laboratory, followed by the immediate administration of corticosteroids in the ED. The receiving ICU can then re-evaluate the continuation of corticosteroids once the results of the ACTH test are available.

Alternatively, ED physicians may administer dexamethasone (4–6 mg IV), which will not interfere with future ACTH stimulation testing and serum cortisol determination and will provide the required glucocorticoid effect.

**M. Blood transfusions in septic ED patients**

**Question**

Should packed red blood cells (PRBCs) be administered to ED patients with early septic shock?

**Recommendation**

During acute sepsis resuscitation, patients with normal CVPs (8–12 mm Hg) and low central venous oxygen saturations (< 70%) should receive blood transfusions to target a Hct of > 30%. (Grade B)

**Rationale**

The optimum hemoglobin (Hgb) level in severe sepsis has not been clearly established. The transfusion requirements in critical care (TRICC) trial studied euvolemic critically ill patients and randomized them to transfusions when a patient’s Hgb was less than 70 g/L versus an Hgb of less than 100 g/L.\(^106\) The primary outcome of 30-day mortality was not significantly different between the 2 groups. Subgroup analysis revealed lower mortality rates in patients with a transfusion threshold of Hgb of 70 g/L in patients who were less ill and younger, and similar mortality rates in patients with coronary artery disease (AMI or angina). However, in a subgroup analysis, there was a trend toward increased death in patients with severe ischemic heart disease. As such, a recommended practice in the euvolemic critically ill patient is to transfuse when the Hgb is less than 70 g/L, or when less than 90–100 g/L if there is a history of coronary artery disease.\(^106\)

However, septic shock patients are often hypovolemic and unstable. As such, they differ from the TRICC trial population. In contrast, in the Early Goal Directed Therapy trial, Rivers and colleagues\(^12\) transfused to achieve a Hct of 30% as part of a sepsis resuscitation algorithm. In this algorithm, transfusion of PRBCs was provided to patients with adequate CVPs and mean arterial pressures, but with low central venous oxygen saturations. Substantial reduction in patient mortality was demonstrated with
the use of this algorithm. Unfortunately, the impact of PRBC administration was not independently assessed. EM physicians should ensure that patients have an adequate amount of oxygen-carrying red blood cells, and if a patient’s central venous oxygen saturation is less than 70%, PRBCs should be administered to achieve a Hct of greater than 30%.

**N. Mechanical ventilation in septic ED patients requiring intubation**

**Question**

1. How should patients with sepsis and respiratory failure be ventilated?

**Recommendation**

Assist control ventilation (ACV) may be used as the initial ventilation mode with an initial FiO₂ of 1.0, positive end expiratory pressure (PEEP) of 5–15 cm H₂O; a rate of 15–25 breaths per minute (bpm). (Grade D)

The goal is to achieve arterial oxygen saturation greater than 90%; a pH greater than 7.25; and a FiO₂ of 0.6 or less. (Grade B)

**Rationale**

The goal of mechanical ventilation is to maintain tissue oxygenation and avoid severe acidosis, while minimizing additional lung damage and the progression to multisystem organ failure. During acute resuscitation, tissue oxygen delivery and the avoidance of severe acidosis (pH < 7.25) are initial physiological goals of ventilation, despite transient high airway pressures and tidal volumes. However, repetitive alveoli collapse and opening is thought to propagate lung injury, and therefore strategies should be employed to minimize barotrauma and volutrauma after patient stabilization.

Although few trials have demonstrated the optimal mode of ventilation, ACV is commonly utilized after emergent airway control as patients are often given sedatives and neuromuscular blockers in the ED. Ventilator settings, such as FiO₂, respiratory rate and PEEP, must be individualized. As such, many patients may be started on ACV, at a FiO₂ of 1.0, PEEP of 5–15 cm H₂O; a rate of 15–25 beats/min initially. Overall, the primary goal is to achieve arterial oxygen saturation greater than 90% and pH less than 7.25. Secondary goals include FiO₂ of 0.6 or less, peak airway pressures of less than 40 cm H₂O and plateau pressures of less than 30 cm H₂O.

Other modes of ventilation, such as pressure support ventilation, may be utilized in more stable patients.

**Question**

2. What ventilator parameters are important to minimize additional lung injury?

**Recommendation**

Patients should receive tidal volumes of 6 mL/kg of ideal body weight and have end-inspiratory plateau pressures of less than 30 cm H₂O to minimize additional lung dysfunction. (Grade B)

**Rationale**

Acute lung injury can result from direct lung injury (e.g., pneumonia, aspiration) or via a systemic response to nonpulmonary disease (e.g., severe nonpulmonary sepsis). Optimal mechanical ventilation may be “protective,” as it minimizes additional ventilator-induced lung injury.

Further lung damage can result from regional overdistention related to high ventilatory volumes (volutrauma), high pressures (barotrauma) and cyclic opening and closing of alveoli with resultant shear injury. A multicentre trial that gradually decreased tidal volumes to 6 mL/kg of ideal body weight (as opposed to 10–12 mL/kg) and aimed for end-inspiratory plateau pressures of less than 30 cm H₂O, showed a 22% decrease in mortality. Decreasing ventilatory volume means permitting moderate hypercapnea. However, “permissive hypercapnea” is safe, providing severe acidosis is avoided, and as long as the patient can hemodynamically tolerate the sedatives required to overcome their respiratory drive. In contrast to initiating this approach immediately, reducing tidal volume over several hours minimizes the hemodynamic consequences of acute hypercapnea (acidosis, vasodilation, increased right ventricular pressure vasodilatation). Sodium bicarbonate, by bolus or infusion, may be required if acidosis becomes pronounced.

**O. Serum glucose control in the ED**

**Question**

Should serum glucose be manipulated in septic ED patients?

**Recommendation**

Both hyperglycemia and hypoglycemia should be avoided in acute sepsis. Blood glucose should be maintained within normal range (4–8 mmol/L) in septic ED patients. (Grade D)

**Rationale**

A single-centre European study of 1600 surgical ICU patients
compared the use of an insulin infusion to maintain blood glucose between 4.4–6.1 mmol/L (“tight glycemic control”) with 10–12 mmol/L (“conventional glucose control”). Tight glycemic control was associated with improved mortality and morbidity. However, questions have remained about its generalizability of tight glucose control for patients with early sepsis, as this study was in postoperative surgical patient. Further evaluation of medical ICU patients also found a morbidity benefit but no overall mortality benefit with tight glycemic control. Unfortunately, this study demonstrated a substantial incidence of hypoglycemia associated with tight glucose control.

As such, it is prudent to avoid both hyperglycemia and hypoglycemia in the acute treatment of sepsis. Studies have not focused on ED patients, and several multicentre ICU trials are still underway to confirm the benefit of tight glucose control. However, without invasive and frequent monitoring, attempts at such tight control in the ED may put the patient at risk for dangerous hypoglycemia.

According to the available data, a euglycemic range of 4–8 mmol/L appears reasonable in ED patient.

In patients who are critically ill, reduced blood flow to skin and muscles may limit insulin uptake with subcutaneous or intramuscular routes. IV insulin should be used for glucose control in the ED.

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**Other support provided by:** Marlene Murphy, Administration Assistant, Department of Medicine, Division of Critical Care Medicine, Queen Elizabeth II Health Sciences Centre, Capital District Health Authority, Dalhousie University; David Urquart, Database Consultant, Department of Emergency Medicine, Dalhousie University.

**Competing interests:** None declared.

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