Compilations of key articles and guidelines in a particular clinical practice area are useful not only to clinicians who practice in that area, but to all clinicians. We compiled pertinent articles and guidelines pertaining to drug therapy in the intensive care setting from the perspective of actively practicing critical care pharmacists. This document differs from the original 2002 version in that a broader assembly of intensive care practitioners was involved in the compilation.

Key Words: pharmacology, intensive care unit, ICU, critical care, key articles, guidelines.

Association for the Surgery of Trauma [EAST] Ad Hoc Committee on Practice Management Guideline Development. Utilizing evidence-based outcome measures to develop practice management guidelines: a primer; available from http://www.east.org/tpg.html). This classification system is as follows: class I, randomized controlled trials; class II, prospective trials and retrospective studies with reliable data (e.g., case-control studies); and class III, retrospective studies and expert opinion.

The information in this compilation should be particularly useful for trainees and relatively new practitioners in the critical care setting. However, we hope it also might serve as a useful template for experienced clinicians who have contemplated a similar undertaking in their practice areas.

The development process began with a preliminary discussion of a potential revision of the original article by several Critical Care PRN members. These members were subsequently sent a copy of the original article and asked for input regarding possible format changes (e.g., disease- vs drug-based approach) and choice of sections for authorship. Additional authors from the Critical Care PRN were then invited to participate based on their recognized areas of expertise in sections not chosen by members of the initial group. Once written, the sections were compiled and reviewed by all authors to help ensure appropriate article selections and summaries. We could not include every published paper we considered important to critical care pharmacotherapy; however, we compiled what we believed were the most representative articles addressing the selected critical care topics.

As in the original version, this bibliography is divided into three general sections. The first focuses on landmark and contemporary pivotal articles and guidelines pertaining to pharmacotherapy in the ICU setting. When these were not available in a specific clinical area, comprehensive reviews or articles were chosen that have generated substantial discussion among ICU practitioners. The second section focuses on literature pertaining to the economic justification of ICU pharmacy services (both supporting and opposing) or, more specifically, the justification of pharmacists in the critical care setting. The final section focuses on articles pertaining to medication errors and adverse drug events in the ICU. Many of the articles pertaining to adverse drug events in the institutional setting apply to the ICU, but we chose articles specific to the ICU.

Pharmacotherapy in the Intensive Care Unit
Acid-Base Disorders


This article reviews the pharmacologic buffering therapy available to treat metabolic acidosis in the ICU. The agents reviewed include sodium bicarbonate, Carbicarb (an equimolar solution of sodium bicarbonate and sodium carbonate), tromethamine, dichloroacetate, thiamine, and volume expanders. Evidence for the clinical efficacy of these buffering agents and correction of arterial pH are also reviewed.


The practice of critical care requires an advanced understanding of acid-base physiology. This article reviews three variables that regulate blood pH: carbon dioxide, relative electrolyte concentrations, and total weak acid concentrations. It provides pathophysiologic mechanisms by organ system and differential diagnoses of metabolic acid-base disorders.


This is the first of a two-part review article that focuses on consequences of severe acidemia. It reviews the management of metabolic acidosis (lactic acidosis, diabetic ketoacidosis, alcoholic ketoacidosis, methanol and ethylene glycol intoxications, aspirin intoxication, toluene exposure, bicarbonate loss, renal failure and dilutional acidosis) and respiratory acidosis.


Part II of the two-part article reviews the adverse cardiovascular, respiratory, metabolic and cerebral consequences of severe alkalemia. Management strategies for metabolic alkalosis, respiratory alkalosis, pseudorespiratory alkalosis, and mixed alkaloses are discussed.

Acute Respiratory Distress Syndrome

In recent years, advances have been made in the treatment and general understanding of acute respiratory distress syndrome. This comprehensive review article addresses the definitions, epidemiology, pathogenesis, and treatment of the syndrome. The authors concluded that future trials targeting new ventilatory strategies or pharmacotherapy may further demonstrate a reduction in mortality.


Ketoconazole possesses antiinflammatory properties that may be beneficial for the treatment or prevention of early acute lung injury or acute respiratory distress syndrome. In this randomized controlled trial, no significant differences in survival, duration of mechanical ventilation, or lung function were found between ketoconazole 400 mg/day (117 patients) and placebo (117 patients). This article is important because it provided firm evidence that ketoconazole is not useful for treatment of acute respiratory distress syndrome. (Class I)

Alcohol Withdrawal


This randomized, double-blind, controlled trial evaluated the potential benefits of individualized therapy compared with fixed-schedule dosing of benzodiazepine for prevention of alcohol withdrawal. Benzodiazepine therapy was administered either as a fixed dose or based on the patient’s score on the revised Clinical Institute Withdrawal Assessment–Alcohol, using a placebo-controlled, double-dummy design. Symptom-triggered therapy resulted in a significant reduction in dosage and duration of therapy compared with the fixed-dosage regimen. The authors concluded that symptom-triggered therapy can shorten detoxification time and reduce unnecessary drug administration without compromising safety. The ability to extrapolate these findings to critically ill patients experiencing alcohol withdrawal is unknown. (Class I)

D’Onofrio G, Rathlev NK, Ulrich AS, Fish SS,


This prospective, randomized, double-blind, placebo-controlled trial evaluated whether lorazepam prevented recurrent seizures in chronic alcoholics. One hundred eighty-six patients with a witnessed generalized seizure, known to have had a drink within 72 hours of seizure onset, received either lorazepam or placebo in addition to routine care. Patients were observed for 6 hours for evidence of recurrent alcohol-related seizures. The risk of subsequent seizures was significantly higher in the placebo group (odds ratio [OR] 10.4, 95% confidence interval [CI] 3.6–30.2). Based on these findings, benzodiazepines appear to be effective agents for the prevention of recurrent alcohol withdrawal seizures. (Class I)


This article presents an evidence-based guideline for the pharmacologic management of alcohol withdrawal based on a meta-analysis of the literature. Six prospective, controlled trials, involving five different agents, demonstrated that benzodiazepines are the most efficacious drug therapy for reducing signs and symptoms of alcohol withdrawal. The superiority of one benzodiazepine over another was not definitively demonstrated. The use of a structured assessment, such as the revised Clinical Institute Withdrawal Assessment–Alcohol, is recommended for evaluating and monitoring patients at risk for alcohol withdrawal. Treatment guidelines and a discussion regarding the efficacy of potential adjunctive options are provided. This guideline did not specifically address the management of alcohol withdrawal in critically ill patients.

Cardiovascular

The following compilations of key articles and guidelines pertaining to arrhythmias, acute coronary syndromes, and hypertension have been covered in previous issues of this journal:

Key articles and guidelines in pharmacotherapeutic management of arrhythmias. Pharmacotherapy 2004;24:248–79.


Arrhythmias


Atrial arrhythmias are common in critically ill patients. In this prospective, randomized trial, two different dosage regimens of amiodarone were compared with diltiazem in 60 patients with new-onset atrial tachyarrhythmias (95% had atrial fibrillation). At least 50% of patients in all three groups achieved more than a 30% rate reduction within 4 hours (p=NS). Over a 24-hour period, patients receiving diltiazem demonstrated a significantly greater reduction in heart rate than those in either amiodarone group (p<0.01). However, frequency of hypotension was higher in the diltiazem group (p<0.01), often requiring withdrawal of treatment. The authors concluded that both drugs provided effective rate control in patients with atrial tachyarrhythmias; however, amiodarone may be better tolerated in hemodynamically unstable patients. (Class I)


The authors cite drug-induced acute pulmonary toxicity in critically ill patients as a reason to reevaluate the use of amiodarone in the ICU. This article reviews dosing, pharmaco-dynamics, pharmacokinetics, drug interactions, efficacy, and toxicity of amiodarone. The authors concluded that there are alternatives to amiodarone for many patients, and caution should be exercised when the drug is used in the treatment of critically ill patients.

Heart Failure


The Vasodilation in the Management of Acute Congestive Heart Failure (VMAC) trial enrolled 489 patients with New York Heart Association class III or IV heart failure at 55 medical centers in the United States. The primary end points were changes in pulmonary capillary wedge pressure (PCWP) and patient-reported dyspnea scores at 3 hours. Secondary outcome measures were PCWP changes at 24 hours, dyspnea scores, global clinical status, and safety. Patients were initially stratified based on presence or absence of a right heart catheter. At the end of 3 hours, the mean change in PCWP was -5.8 mm Hg for patients receiving nesiritide, -3.8 mm Hg for those receiving nitroglycerin, and -2 mm Hg for those receiving placebo (p<0.05 for comparison of nesiritide with nitroglycerin and placebo). Dyspnea scores were significantly improved when nesiritide was compared with placebo but was no different when compared with nitroglycerin. Global clinical status was similar in all groups (p=NS). After 24 hours, PCWP changed by -8.2 mm Hg in the nesiritide group compared with -6.3 mm Hg in the nitroglycerin group (p=0.04). Dyspnea and global clinical status were not significantly different. Adverse events were more common in the nitroglycerin group than the nesiritide group (p<0.001). Headache and pain were reported most often. Rates of hypotension, ischemic events, and arrhythmias were similar between the groups. However, duration of hypotension was significantly longer with nesiritide (2.2 hrs) than nitroglycerin (0.7 hrs) (p=0.002). This trial provides additional evidence for the conclusions of a previous trial (see Colucci WS, Elkayam U, Horton DP, et al. Intravenous nesiritide, a natriuretic peptide, in the treatment of decompensated congestive heart failure. N Engl J Med 2000;343:246–53) that nesiritide is a useful addition to standard therapy in the treatment of patients with decompensated heart failure. (Class I)


These guidelines present a graded, evidence-based approach to managing heart failure. They provide an overview of public health concerns, pathophysiology, and therapy. The guidelines consider angiotensin-converting enzyme inhibitors and diuretics fundamental aspects of heart failure treatment. Graded recommendations are made.
for adding adrenergic receptor blockers, digoxin, anticoagulation and antiplatelet agents, angiotensin receptor blockers, antiarrhythmics, and aldosterone antagonists. Treatment of myocarditis is also reviewed.

Hypertension


This article reviews the underlying pathophysiology, hemodynamics, potential complications, and treatment options for acute postoperative hypertension. The authors present the pharmacology, clinical response, and supporting evidence for the vasodilators and β-blockers commonly used in the management of acute postoperative hypertension, along with the significant limitations of the current literature.

Fluids


This multicenter, randomized, double-blind comparison of 4% albumin and normal saline for resuscitation of patients admitted to the ICU enrolled 6997 patients; 28-day mortality was the primary end point. No statistically significant differences were observed in death (relative risk [RR] 0.99, 95% CI 0.91–1.09) or any of the secondary end points under study. (Class I)


This was a double-blind, randomized, controlled trial involving 229 hypotensive patients (systolic blood pressure < 100 mm Hg) who experienced traumatic brain injury (Glasgow Coma Scale score < 9). In addition to conventional resuscitation procedures employed by paramedics, patients were randomized to receive rapid intravenous infusions (250 ml) of either 7.5% sodium chloride or Ringer’s lactate solution. Patients in both groups were given additional crystalloid or colloid fluids based on protocol. The major outcome measure was the extended Glasgow Outcome Score at 6 months. Survival rates were similar with 7.5% sodium chloride and Ringer’s lactate (55% and 50%, respectively, at 6 mo, p=0.23). Similarly, no significant difference was noted between the groups based on neurologic function (p=0.96). The use of the Glasgow Outcome Score as a primary outcome measure in this study is somewhat of a concern in that it is more reflective of unfavorable rather than favorable outcomes. However, as mentioned by the investigators in their response to this criticism (JAMA 2004;291:2944–5), a much larger sample would have been needed to investigate the effects of the solutions based on a favorable outcome measure. (Class I)


Renal impairment, as a result of cytokine release and decreased arterial blood volume, is common in patients with spontaneous bacterial peritonitis. Albumin may prevent renal dysfunction through its effects on volume expansion and possibly other mechanisms. In this study, 126 patients with cirrhosis and spontaneous bacterial peritonitis were randomized to cefotaxime or cefotaxime plus albumin. In the control group, 33% of patients developed renal impairment, versus 10% in the albumin group (p=0.002). Mortality in the hospital and at 3 months was significantly lower in albumin-treated patients than in control patients (p=0.01 and p=0.03, respectively). This trial is important because it is one of the few well-controlled trials demonstrating a positive benefit of albumin beyond its effects on surrogate end points. (Class I)


This review of 17 randomized trials compared isotonic crystalloids with colloids for volume resuscitation in 814 adult patients. No differences were noted in mortality, pulmonary edema, or length of hospital stay. However, a subgroup analysis revealed decreased mortality with crystalloid administration in patients who experienced trauma (RR 0.39, 95% CI 0.17–0.89). The authors concluded that more research is necessary to resolve the ongoing controversy regarding the most appropriate and effective method for fluid resuscitation.

Cochrane Injuries Group Albumin Reviewers. Human albumin administration in critically ill

In this systematic review of 30 randomized, controlled trials involving 1419 critically ill patients, albumin administration was associated with one death for every 17 patients treated for hypovolemia, burns, or hypoalbuminemia. Although a significant difference in mortality after albumin administration was not found in each of the categories, the pooled relative risk of albumin-associated death was significant (RR 1.68, 95% CI 1.26–2.23). This review, as well as an earlier review in the same journal (Schiehout G, Roberts I. Fluid resuscitation with colloid or crystalloid solutions in critically ill patients: a systematic review of randomised trials. BMJ 1998;316:961–4), fueled the colloid-crystalloid debate by pushing the limits of meta-analytic technique.


Previous studies in animal models have suggested that the volume of intravenous fluid resuscitation may contribute to intraoperative bleeding. In this prospective study, trauma patients with hypotension were randomized to immediate (309 patients) or delayed (289 patients) resuscitation. The primary objective was to determine if the rate of survival would improve when intravenous fluid resuscitation was withheld until the time of surgery. The overall rate of survival was significant in the delayed versus immediate resuscitation group (70% vs 62%, p=0.04). Postoperative complications (acute respiratory distress syndrome, sepsis syndrome, acute renal failure, coagulopathy, wound infection, and pneumonia) also tended to occur less often when fluid administration was delayed, although the difference was not significant. (Class I)

Endocrine Conditions


This prospective study assessed baseline serum total cortisol, cosyntropin-stimulated serum total cortisol, aldosterone, and free cortisol concentrations in 60 critically ill patients and 33 healthy volunteers. Critically ill patients were stratified into two groups based on serum albumin concentration. Baseline and cosyntropin-stimulated total cortisol concentrations were significantly lower in patients whose albumin concentrations were 2.5 g/dl or less. However, average baseline free cortisol concentrations were not different between the two groups of critically ill patients. Free cortisol concentrations were more than 8 times higher in the critically ill patients than in the healthy volunteers. The authors concluded that caution should be used in interpreting total cortisol concentrations in critically ill patients with hypoalbuminemia, and that measuring serum free cortisol concentrations may prevent unnecessary use of glucocorticoids. (Class II)


This is a secondary analysis of a previously published report demonstrating the efficacy of intensive insulin therapy in reducing morbidity and mortality among critically ill patients. The authors focused their assessment on the feasibility and safety of intensive insulin therapy, identification of factors that determine insulin requirements, and determination of the impact of glycemic control versus the amount of exogenous insulin infused on morbidity and mortality measures. The analysis demonstrated that it is safe and feasible to maintain blood glucose levels of 80–110 mg/dl using a titration algorithm. Independent factors influencing insulin requirements were body mass index, history of diabetes mellitus, reason for ICU admission, presence of hyperglycemia on admission, caloric intake, and duration of ICU stay. Multivariate logistic regression analysis revealed that lower blood glucose level rather than insulin dosage was related to reduced morbidity and mortality. Among patients whose ICU stay exceeded 5 days, post hoc analysis revealed a gradual decrease in risk of death with decreasing blood glucose level. Patients whose blood glucose was maintained at intermediate levels of 110–150 mg/dl had worse outcomes than those whose levels were normal. (Class II)


This review eloquently summarizes the physiology of corticosteroid response to acute illness by contrasting both the appropriate and
inappropriate hypothalamic-pituitary-adrenal axis response during acute illness. The authors describe the clinical features of hypoadrenalism in patients with acute illness. They also provide a useful algorithm for evaluating adrenal corticosteroid function in critically ill patients based on random cortisol levels and/or the corticotropin stimulation test. Recommendations are provided for steroid replacement during acute illness in patients with adrenal insufficiency, including those who previously received corticosteroids.


This prospective, randomized, controlled trial involving 1548 mechanically ventilated surgical patients with hyperglycemia evaluated the effects of intensive insulin therapy or conventional treatment. Goal blood glucose levels were 80–110 mg/dl with intensive insulin therapy and 180–200 mg/dl with conventional treatment. At 12 months, mortality was significantly reduced in the intensive insulin therapy group compared with the conventional treatment group (4.6% vs 8%, p<0.04). The benefits of intensive insulin therapy were most noteworthy in the patients whose ICU stay exceeded 5 days; mortality was 10.6% in that intensive insulin therapy group compared with 20.2% in the conventional treatment group (p<0.005). This trial also demonstrated significant reductions in morbidity (e.g., bloodstream infections, acute renal failure requiring dialysis, number of blood transfusions, critical-illness polyneuropathy, duration of mechanical ventilation, and length of ICU stay) with the use of intensive insulin therapy. (Class I)


This comprehensive review discusses the pathogenesis, differential diagnosis, and treatment strategies for acute and chronic water deficits. Desmopressin acetate is the preferred agent for treatment of symptomatic central diabetes insipidus, whereas treatment of nephrogenic diabetes insipidus centers on the resolution of underlying conditions. The authors identified noncompliance, medical emergency, and iatrogenic inadvertence as the most common causes of unplanned treatment withdrawals.

Fever


This review summarizes the literature regarding whether or not a fever should be treated and the effectiveness of various interventions for treating fever in critically ill patients. Although the author does not use the more traditional classifications of levels of evidence, in which level 1 is the highest, the classification used is provided and readily understandable. The review highlights the tremendous lack of data evaluating various strategies (e.g., physical cooling vs antipyretics) used to treat fever.


Fever in intensive care patients consumes substantial financial and medical resources. An interdisciplinary team with members from the areas of critical care, infectious diseases, and surgery developed consensus guidelines to aid in the rational, cost-conscious evaluation of a new fever. The overall aim of these evidence-based guidelines, endorsed by the Society of Critical Care Medicine and the Infectious Diseases Society of America, is to minimize unnecessary testing and facilitate prompt implementation of appropriate therapy.

Gastrointestinal Conditions

Intestinal Transit


This prospective, randomized, double-blind trial evaluated the efficacy of enteral naloxone (8 mg every 6 hrs) versus placebo in reducing gastric tube reflux and the frequency of pneumonia in 84 mechanically ventilated patients. All patients were receiving continuous infusion of fentanyl for analgesia, and the dosage requirements were not significantly different between the two groups. The median daily gastric tube reflux volumes were significantly lower in the naloxone treatment group than in the placebo group (34 vs 129 ml, p=0.03). The frequency of pneumonia was also significantly lower in the naloxone versus the placebo group (34.2% vs 55.8%, p=0.04). The authors concluded that oral naloxone might be a simple and possibly preventive strategy to reduce gastric tube reflux and the frequency of pneumonia in
fentanyl-treated, mechanically ventilated patients. (Class I)


Delayed gastric emptying is common among critically ill patients, occurring in up to 60% of those who are mechanically ventilated. Gastric hypomotility may lead to impaired nutrient delivery, high gastric residuals, and risk of aspiration. This review summarizes the data regarding the efficacy of prokinetic agents (metoclopramide, erythromycin, and cisapride) in improving gastric emptying in critically ill patients. Eight of the 10 studies reviewed demonstrated a positive effect. However, four of these studies evaluated only cisapride, which has been removed from the United States market. Although evidence supports the efficacy of prokinetic agents (erythromycin and metoclopramide) for promoting gastric emptying, data demonstrating a positive impact on patient outcome (e.g., reduced frequency of pneumonia or mortality) are lacking.


In this double-blind, randomized, placebo-controlled trial of 30 ventilated patients with multiple organ failure, 11 of 13 responded to neostigmine treatment for critical illness–related colonic ileus; resolution of symptoms was not evident in any of the patients in the placebo group (11 neostigmine-treated patients vs 0 placebo patients, p<0.001). Nineteen (79%) of 24 neostigmine-treated patients defecated, including nonresponders who were switched from the placebo group. Continuous infusion of neostigmine (0.4–0.8 mg/hr for 24 hrs) appeared to be a well-tolerated therapy in this small sample of critically ill, mechanically ventilated patients with a colonic ileus. (Class I)

Stress Ulcer Prophylaxis


This study uses three different strategies to estimate mortality and length of stay in the ICU attributable to clinically important upper gastrointestinal bleeding. Data from 1666 mechanically ventilated patients from two multicenter databases were used to perform the analyses. Of the 1666 patients, 59 developed clinically important gastrointestinal bleeding. Risk of death was increased in patients with bleeding, regardless of analysis strategy used; relative risk ranged from 1.8 (95% CI 1.1–2.9) to 4.1 (95% CI 2.6–6.5). The median length of ICU stay attributable to clinically important bleeding was 3.8–7.9 days depending on the model employed. Overall, this study demonstrates that although the frequency of clinically important gastrointestinal bleeding may be low, the patient’s condition is associated with important attributable morbidity, such as a long ICU stay (4–8 days), and mortality. (Class II)


Numerous articles have been published on stress ulcer prophylaxis. This comprehensive, evidence-based guideline reviews the frequency of bleeding and the efficacy of prophylaxis in intensive care patients and special populations. Although this 1999 review lacks the more recent findings, it summarizes the epidemiology and populations at risk for stress ulcers. It also evaluates the evidence regarding treatment options available at the time. A more recent article (Allen ME, Kopp BJ, Erstad BL. Am J Health-Syst Pharm 2004;61:588–96) critically reviews more recently published data on therapeutic approaches to stress ulcer prophylaxis.


Two agents used for stress ulcer prophylaxis, ranitidine and sucralfate, were compared in 1200 mechanically ventilated, critically ill patients for prevention of clinically important upper gastrointestinal bleeding. This landmark, multicenter, randomized, double-blind, placebo-controlled study demonstrated a lower bleeding frequency in the ranitidine group than in the sucralfate group (RR 0.44, 95% CI 0.21–0.92).
No difference between the two groups was noted for rate of ventilator-associated pneumonia, length of ICU stay, or survival. This study demonstrates the superiority of ranitidine (50 mg given intravenously every 8 hrs) over sucralfate (1 g administered by nasogastric tube every 6 hrs) for prevention of clinically important upper gastrointestinal bleeding. (Class I)


This prospective, multicenter, observational study evaluated potential risk factors for stress ulceration in 2252 critically ill patients. Clinically important gastrointestinal bleeding occurred in 1.5% of patients. The two major independent risk factors for bleeding identified by multivariate analyses were respiratory failure (OR 15.6), and coagulopathy (OR 4.3). This study clearly highlights the importance of respiratory failure and coagulopathy as two major risk factors for stress-related mucosal bleeding. However, because the study involved only a critically ill patient population, these should not be considered the only risk factors for stress-related mucosal bleeding. Specifically, earlier literature suggests that patients with head injury, burns, multiple trauma, sepsis, or shock may also be at risk. However, few patients with these conditions were included in the trial. (Class II)

Nonvariceal Upper Gastrointestinal Bleeding


Consensus guidelines for the management of nonvariceal upper gastrointestinal bleeding were developed by a multidisciplinary panel representing 11 national societies. The guidelines recommend that institution-specific protocols be developed. Initial management should include stratification of patients into high and low risk categories for rebleeding and death on the basis of clinical, endoscopic, and prognostic factors. Early endoscopy is warranted to facilitate diagnosis and risk stratification, and to provide hemothostasis in patients with stigmata suggesting high risk of rebleeding. A combination of injection and thermal coagulation provides superior endoscopic hemothostasis in patients with high-risk stigmata compared with either treatment alone. High-dose intravenous therapy with a proton pump inhibitor is recommended for patients who have undergone endoscopic hemothostasis. Each recommendation was graded according to the level and strength of available evidence.


Earlier trials demonstrated that omeprazole given as a continuous infusion is effective in preventing peptic ulcer rebleeding in patients at high risk of rebleeding (i.e., nonbleeding visible vessel or bleeding ulcers) who have undergone endoscopic hemostasis. This single-blind, randomized trial assessed whether combination therapy (endoscopy plus a proton pump inhibitor) was superior to omeprazole alone in preventing peptic ulcer rebleeding in patients with nonbleeding visible vessels or adherent clots. The probability of recurrent bleeding within 30 days was significantly less in the group receiving combination therapy (1.1%) than in the group receiving omeprazole alone (11.6%; p=0.009). Previously published data demonstrating the efficacy of omeprazole after endoscopic hemostasis, in concert with the results of this study, emphasize that dual-modality therapy (endoscopic hemostasis plus intravenous omeprazole) is the most effective strategy for reducing the risk of rebleeding in high-risk patients. (Class I)


This meta-analysis was conducted to assess the efficacy of histamine2 (H2)-receptor antagonists in the management of peptic ulcer bleeding. Pooling data from randomized placebo-controlled trials, the authors demonstrated that H2-receptor antagonists do not significantly reduce rebleeding, surgery, or death in patients with bleeding duodenal ulcers. Significant, however minor, reductions were observed in rebleeding, surgery, and death in patients with bleeding gastric ulcers (absolute risk reduction 7.2%, 6.7%, and 3.2%, respectively). Based on these data, intravenous H2-receptor antagonists should be considered of no value in patients with bleeding duodenal ulcers and of limited value in patients with bleeding gastric ulcers.

This meta-analysis evaluated whether proton pump inhibitors were more effective than H₂-receptor antagonists in patients with bleeding peptic ulcers. Using data from 11 international comparative, randomized trials, persistent or recurrent bleeding was noted in 6.7% (95% CI 4.9–8.6%) of patients receiving proton pump inhibitors and in 13.4% (95% CI 10.8–16%) of patients treated with H₂-receptor antagonists. Although this analysis demonstrates that proton pump inhibitors are more effective than H₂-receptor antagonists in preventing persistent or recurrent peptic ulcer bleeding, no significant difference was seen in mortality or need for surgery.


This landmark, single-center, randomized, double-blind, placebo-controlled trial assessed the efficacy of intravenous omeprazole in preventing peptic ulcer rebleeding after endoscopic hemostasis. Patients with actively bleeding ulcers or ulcers with nonbleeding visible vessels received endoscopic hemostasis with epinephrine injection followed by thermo-coagulation. Then, 240 patients were randomized in a double-blind fashion to receive an intravenous bolus dose of omeprazole 80 mg followed by 8 mg/hour continuous infusion, or placebo for 72 hours. Subsequently, all patients received oral omeprazole 20 mg/day for 8 weeks. The frequency of rebleeding within the first 30 days was significantly reduced in the omeprazole group (6.7%) compared with the placebo group (22.5%). Most episodes of bleeding occurred within the first 3 days. This well-designed trial demonstrates the efficacy of omeprazole given as a high-dose continuous infusion for prevention of peptic ulcer rebleeding after endoscopic hemostasis in patients at high risk of rebleeding. (Class I)

Hematology

Antithrombotics


This report from the American College of Chest Physicians provides an extensive evidence-based review of the management of thromboembolic disorders.

Blood Conservation and Transfusion


This prospective, multicenter, observational, cohort study of 4892 patients, conducted in 2000–2001, serves as an important historic control to compare current patterns of blood use. The patients’ mean ± SD hemoglobin level was 11 ± 2.4 g/dl, with a progressive decrease throughout the ICU stay. Overall, 44% of patients received at least one transfusion of red blood cells. Most transfusions were given in the first week, and thereafter at a rate of 1–2 units/week. Mean ± SD time to first transfusion was 2.3 ± 3.7 days (median 1 day). Transfusion of red blood cells was associated with worsened patient outcomes. These data were similar to Western European ICU data collected in 1999, when hemoglobin at baseline, hemoglobin at the time of transfusion, and association between mortality and transfusion were evaluated (Vincent JL, Baron J-F, Reinhart K, et al. Anemia and blood transfusion in critically ill patients. JAMA 2002;288:1499–507). (Class II)


This literature review focuses on the use of transfusions for acute management of anemia in critically ill patients. The risks and benefits of red blood cell transfusion and the use of blood substitutes are reviewed. Despite measures to reduce blood wastage, many patients develop anemia of critical illness, characterized by low red blood cell production despite normal-to-high concentrations of erythropoietin, and in some cases inadequate erythropoietin production for the degree of anemia. Exogenous erythropoietin has raised red blood cell production and corrected anemia in a variety of patient populations. The authors make recommendations for therapy based on the limited data and expert opinion available.

Corwin HL, Gettinger A, Pearl RG, et al.

This randomized controlled trial compared erythropoietin 40,000 U/week with placebo in a selected medical-surgical ICU population. Transfusion independence was the primary end point. The erythropoietin-treated patients were less likely than placebo patients to receive a red blood cell transfusion during the 28-day study, but their hemoglobin increase was greater than that of placebo patients. No significant adverse effects were reported. The regimen and end points were different from those used in an earlier pilot study (Corwin HL, Gettinger A, Rodriguez RM, et al. Efficacy of recombinant human erythropoietin in the critically ill patient. Crit Care Med 1999;27:2346–50). This trial, performed before another study (Hébert PC, Wells G, Blajchman MA, et al. A multicenter, randomized, controlled clinical trial of transfusion requirements in critical care. N Engl J Med 1999;340:409–17), compared restrictive versus liberal transfusion, resulted in a transfusion trigger higher than currently advocated. (Class I)


The authors searched their clinical database for information regarding nosocomial infection in 1717 patients in their medical-surgical-trauma ICU for admissions from October 1998–August 2000. Risk of infection was calculated based on the entire cohort—patients receiving and not receiving red blood cell transfusion. The overall nosocomial infection rate was 5.94%, and the rates were significantly different between the patients who received (15.3%) and did not receive (2.9%) transfusion (p<0.005). This difference persisted after adjusting for severity of illness with scores from the Mortality Prediction Model. In addition, an association was noted between the number of units administered and risk of infection. Transfused patients had longer ICU and hospital stays (p<0.0005). The risk of nosocomial infection, although not proven to be the direct result of red blood cell transfusion, may be the result of transfusion-related immunosuppression. Similar results have been documented in other populations, such as trauma patients (Claridge JA, Sawyer RG, Schulman AM, et al. Blood transfusions correlate with infections in trauma patients in a dose-dependent manner. Am Surg 2002;68:566–72). (Class II)


This article, the first of a two-part series, extensively reviews the risks of blood transfusion, such as human immunodeficiency virus, hepatitis B and C, and other hepatitis viruses. Other risks, such as hemolytic reactions, red cell bacterial contamination, transfusion-related lung injury, and immunomodulation are also reviewed. The indications for transfusion reflect the thinking at the time, although these have been altered by subsequent reports as described below.


These investigators randomized patients to either a liberal transfusion strategy designed to maintain a hemoglobin level of 10–12 g/dl or a restrictive strategy designed to maintain a hemoglobin level of 7–9 g/dl. More red blood cell transfusions were given to the liberal group, and although the primary outcome of all-cause mortality was not different between the groups, the in-hospital mortality and multiple-organ dysfunction score were significantly higher in the liberal group. These data indicate that a transfusion threshold as low as 7 g/dl was at least as effective and possibly superior to a more liberal transfusion strategy. A trend toward a lower survival was noted in a subset of patients with cardiovascular disease from the restrictive transfusion group. This led the authors to suggest that patients with unstable coronary ischemic syndromes may not benefit from a restrictive transfusion strategy (Hébert PC, Yetisir E, Martin C, et al. Is a low transfusion threshold safe in critically ill patients with cardiovascular disease? Crit Care Med 2001;29:227–34). (Class I)

Infection

Bacterial

This report updates, expands, and replaces the previously published Guidelines for Prevention of Nosocomial Pneumonia. The new guidelines are designed to reduce the frequency of pneumonia and severe, acute lower respiratory tract infections in acute-care hospitals, other health care settings (e.g., ambulatory and long-term care institutions) and other facilities where health care is provided. The changes in the recommendations to prevent bacterial pneumonia—especially ventilator-associated pneumonia—are directed at endotracheal intubation, water and environmental purification to avoid Legionella contamination, administration of palivizumab to prevent respiratory syncytial virus infection, use of newer antiviral agents for treatment of influenza, and identification of pertussis and lower respiratory tract infections caused by adenovirus and human parainfluenza viruses.


A prospective, randomized, double-blind, clinical trial in 401 patients with ventilator-associated pneumonia conducted in 51 French ICUs found no difference in mortality or recurrent infection in patients treated with antibiotics for 8 days compared with those treated for 15 days. No differences were noted in resolution of fever, leukocyte count, the ratio of partial pressure of arterial oxygen to fraction of inspired oxygen, organ dysfunction, or radiologic scores between the two treatment groups. In a secondary analysis, there were more patients with nonfermenting gram-negative bacilli infections treated with 8 days of antibiotics who had pulmonary infection recurrence. These data demonstrated that for intensive care patients with ventilator-associated pneumonia, there is no clinical advantage in prolonging antimicrobial therapy to 15 days compared with 8 days. (Class I)


This is a retrospective analysis of the empiric treatment of P. aeruginosa bacteremia in 115 episodes over 10 years, from 1988–1998. The work is unique in that 30-day mortality was an end point. Empiric therapy with combination antibiotics was associated with a better 30-day survival rate than empiric monotherapy. No difference was found in 30-day survival with monotherapy versus combination therapy if a definitive treatment was used after receipt of the culture and sensitivity report. These data support the long-held principle that P. aeruginosa bacteremia is better treated with a combination of antipseudomonal antibiotics than with monotherapy, at least until culture and susceptibility data have been received. (Class III)


This meta-analysis reviewed mortality associated with both methicillin-susceptible S. aureus (MSSA) and methicillin-resistant S. aureus (MRSA) bacteremia in 31 studies that included 3963 patients; 65.7% had MSSA, 34.3% MRSA. No significant difference in mortality was found in 24 studies; higher mortality associated with MRSA bacteremia was found in seven studies. When all data were combined, a significant increase in mortality associated with MRSA bacteremia was observed. This increased rate was also observed in a subgroup analysis in which adjustment was made for confounding variables.


This is a post hoc analysis of two randomized, double-blind, multicenter studies of linezolid versus vancomycin for gram-positive nosocomial pneumonia. Data on 1019 patients were combined; 339 patients had documented S. aureus pneumonia, and 160 had documented MRSA pneumonia. Clinical cure rates for MRSA pneumonia were 50% for linezolid-treated patients and 35.5% for vancomycin-treated patients (p<0.01). The 28-day survival rate for patients with MRSA pneumonia was 80% with linezolid versus 63.5% with vancomycin (p=0.03). After adjusting for baseline variables, the difference favoring linezolid remained significant. Although this article is a retrospective analysis of combined data, it is the first to demonstrate a survival advantage for one appropriate antibiotic regimen over another in patients treated for MRSA pneumonia. (Class III)

Antibiotic susceptibility results for 35,790 nonduplicate gram-negative aerobic bacilli recovered from patients from 1994–2000 representing 43 states and the District of Columbia were reported. These results were compared with those from a similar analysis of data for 1990–1993. The activity of most antimicrobial agents decreased 6% or less over the study period. Ciprofloxacin activity decreased from 89% in 1990–1993 to 86% in 1994 to 76% in 2000. The decrease in activity of ciprofloxacin was associated significantly with the increased national use of fluoroquinolone therapy during the study period. Cross-resistance to other broad-spectrum antibiotics—such as gentamicin, ceftazidime, imipenem, and amikacin—was observed in ciprofloxacin-resistant isolates.


This prospective, randomized, double-blind, multicenter trial involved 301 adult patients with bacterial meningitis. Results demonstrated that dexamethasone 10 mg administered just before or along with the first dose of antibiotic (intravenous amoxicillin) and then continued every 6 hours for 4 days was associated with a reduced risk of unfavorable outcome as indicated by the Glasgow Outcome Scale. Dexamethasone significantly affected unfavorable outcome in patients with pneumococcal meningitis but not meningitis due to Neisseria meningitides. The proportion of patients who died was significantly smaller in the dexamethasone group. Dexamethasone did not have a beneficial effect on neurologic sequelae, including hearing loss. Although management of meningitis differs throughout the world, these results support early treatment with dexamethasone in adults with bacterial meningitis. The direct applicability of these data in countries where vancomycin is the first-line therapy remains to be determined. (Class I)


This article presents consensus guidelines for diagnosis and treatment of ventilator-associated pneumonia. Twelve European intensivists were posed 21 questions regarding ventilator-associated pneumonia. Answers were independently provided, and then were reported to and discussed among all participants. The guidelines cover several important topics, including initial empiric antibiotic therapy, monotherapy versus combination antibiotic therapy, combination therapy for P. aeruginosa infection, consideration of MRSA, duration of antibiotic therapy, and deescalation of antibiotic therapy for the management of ventilator-associated pneumonia. Evidence for the recommendations is provided.


The authors present results from a prospective 2-year trial of protocol-driven antibiotic rotation in a surgical ICU. In year 1, current practice was observed; in year 2, quarterly antibiotic rotation was adopted. Patients were stratified for rotation based on diagnosis of either pneumonia, or peritonitis or sepsis. A significant reduction in both gram-positive and gram-negative bacterial resistance occurred. A significant reduction in mortality associated with infection was identified during the rotation period. Length of ICU stay was not affected. (Class III)


This article demonstrates that critically ill patients who receive initial antimicrobial therapy for an infection that does not have activity against the causative pathogen(s) are significantly more likely to die. Using logistic regression analysis, the investigators found that inadequate antimicrobial treatment of infection was the most important independent predictor of mortality among the 2000-patient cohort. The data also show that up to 45.2% of intensive care patients with infection receive inadequate antimicrobial therapy. (Class II)

Pseudomonas aeruginosa bacteremia is associated with a high mortality rate. The aim of this prospective, observational report, which involved 200 patients, was to determine whether in vitro microbiology data translate into improved outcomes. Factors associated with increased mortality were a respiratory portal of entry, neutropenia, and use of monotherapy. Combination therapy, compared with monotherapy, demonstrated improved survival in patients with P. aeruginosa bacteremia (p<0.02). In vitro synergy testing data did not correlate with improved outcomes. (Class III)

Fungal


These guidelines are extensive and detailed evaluations of the treatment of Candida infections. The guidelines followed the U.S. Public Health Service grading system for rating recommendations in clinical guidelines. A detailed description of susceptibility testing methods for all antifungal agents and their interpretations is provided. Recommendations include updated dosing guidelines for amphotericin B, amphotericin lipid-based formulations, and fluconazole. They also incorporate the use of voriconazole and caspofungin in the management of Candida infections. Specific recommendations regarding fungal prophylaxis are provided.


In this randomized, double-blind, multicenter study of 239 patients with systemic or invasive candidiasis, caspofungin produced 73.4% success, compared with 61.7% for amphotericin B, in an intent-to-treat analysis. In a predefined analysis of patients who met prespecified criteria for evaluation, 80.7% of caspofungin-treated patients and 64.0% of amphotericin-treated patients had successful outcomes (p=0.03). Caspofungin showed consistently superior success among stratified subgroups. Drug-related adverse effects were significantly higher in the amphotericin B group. Failure due to toxic effects requiring change in therapy occurred significantly more frequently in the amphotericin group (16.5%) compared with the caspofungin group (2.8%; p=0.03). These data suggest that caspofungin is an effective yet less toxic alternative to amphotericin B for invasive candidiasis. (Class I)


This multicenter, randomized, open-label trial involved 237 nonneutropenic patients with candidemia. Results demonstrated that fluconazole was equivalent to amphotericin B in overall clinical success and survival. Amphotericin B caused significantly greater elevations in serum blood urea nitrogen and creatinine levels, and more hypokalemia, than fluconazole. The mean ± SD doses used in this study (amphotericin B 0.5 ± 0.01 mg/kg/day, total dose 570 ± 40 mg; fluconazole 5.3 ± 0.2 mg/kg/day) are somewhat lower than recommended dosages. These data suggested that fluconazole could be considered for the management of candidemia, which was previously thought treatable only with amphotericin B. (Class I)

Sepsis


These comprehensive guidelines were prepared by critical care and infectious disease experts from 11 international organizations. Recommendations were graded based on a modified Delphi methodology. The guidelines cover initial resuscitation, diagnosis, antibiotic therapy, source control, fluid therapy, vasopressors, inotropic therapy, steroids, drotrecogin alfa (activated), blood product administration, mechanical ventilation, glucose control, renal replacement therapy, bicarbonate therapy, deep vein thrombosis prophylaxis, stress ulcer prophylaxis, considerations for limitations of support, considerations for pediatric patients, and sedation, analgesia, and neuromuscular blockade. This article represents the culmination of years of research in sepsis, septic shock, and critical illness.

This randomized, double-blind, placebo-controlled, parallel-group, multicenter trial involved 300 adults with septic shock. The investigators evaluated the effects of treatment with intravenous hydrocortisone 50 mg every 6 hours and enteral fludrocortisone 50 µg once/day for 7 days on 28-day mortality. Patients were stratified based on the results of a short corticotropin test. For nonresponders, mortality was 63% in the placebo group and 53% in the treatment group (p=0.04). For responders, as well as for all patients regardless of corticotropin test results, there was no significant effect of corticosteroids on survival. For nonresponders, the time to withdrawal of vasopressor support was 3 days shorter with corticosteroid therapy than placebo (p=0.001). No differences in adverse events were noted between treatment groups. These data support corticotropin testing in patients with septic shock and the administration of replacement corticosteroids in patients with adrenal insufficiency. (Class I)


This prospective, randomized trial involved 263 patients with severe sepsis or septic shock in an emergency department. Results showed a significant 16% absolute improvement in in-hospital survival for patients receiving 6 hours of early goal-directed therapy compared with standard therapy before admission to the ICU. Early goal-directed therapy consisted of achieving a central venous pressure of 8–12 mm Hg or greater, mean arterial pressure of 65–90 mm Hg, and central venous oxygen saturation of 70% or greater. A protocol for achieving these goals was used. Urine output of 0.5 ml/kg/hour or greater was also a goal, but there was no specific therapy in the protocol directed toward urine output. Goals were achieved in 99.2% of the early therapy group, compared with 86.1% of the standard therapy group. The authors suggested that goal-directed therapy provided at the earliest stages of severe sepsis and septic shock has significant short- and long-term benefits. These results may affect the quality and timing of resuscitation before enrollment in future trials of sepsis and septic shock. (Class I)


This randomized, double-blind, placebo-controlled, multicenter trial involved 1690 patients with severe sepsis. The authors found that drotrecogin alfa (activated), which is recombinant human activated protein C, infused at 24 µg/kg/hour for 96 hours reduced all-cause 28-day mortality from 30.8% to 24.7% (p<0.005). Reduced relative risk of death was 19.4%. For patients at high risk of death, as predefined by Acute Physiology and Chronic Health Evaluation II score, drotrecogin alfa (activated) produced an absolute reduction in mortality of 13%, from 44% to 31%. The frequency of severe bleeding was higher in the treatment than placebo group (3.5% vs 2.0%, p=0.06). Drotrecogin alfa (activated) was approved by the Food and Drug Administration for treatment of severe sepsis on the basis of this trial. (Class I)

Neurosurgery and Neurology

Spinal Cord Injury


The authors state evidence is insufficient to support treatment standards or guidelines for blood pressure management for patients with acute spinal cord injury. They do, however, recommend avoiding hypotension (systolic blood pressure < 90 mm Hg) and maintaining mean arterial pressure at 85–90 mm Hg for 7 days after injury to improve spinal cord perfusion.


This is one of the most controversial chapters in the guidelines, and the authors recommend that readers review the data and comments to formulate their own opinions regarding treatment options. They state that there is insufficient evidence for treatment standards or guidelines for using corticosteroid therapy. The group recom-
mends that methylprednisolone administered for 24–48 hours can be considered a treatment option for patients with acute spinal cord injury; however, the authors first acknowledge that the evidence suggesting harmful side effects is more consistent than evidence supporting clinical benefit.


In this chapter, prophylaxis of deep vein thrombosis is recommended as a standard of care for patients with acute spinal cord injury. The authors support the use of low-molecular-weight heparins, rotating beds, adjusted-dose heparin, or a combination of these modalities, as well as low-dose heparin in combination with pneumatic compression stockings or electrical stimulation as treatment strategies for prophylaxis. The guidelines state that low-dose heparin therapy alone and oral anticoagulation alone are not recommended. Other suggestions include the administration of various diagnostic tests, 3 months of prophylactic treatment, and vena cava filters for patients not responding to anticoagulation or who have contraindications to anticoagulation and/or mechanical devices.


This article provides a critical evaluation of the results and conclusions of the National Acute Spinal Cord Injury Studies (NASCIS) II and III to determine the influence they should have on practice standards for patients with acute spinal cord injury. The author converted the published data from these trials into the original raw data to further evaluate the effects on outcome. He concluded that although these studies were well designed and well conducted, evidence to support the use of methylprednisolone therapy for patients with acute spinal cord injury is weak and clinically irrelevant. He suggested that administration of methylprednisolone should be considered experimental, and the 48-hour treatment regimen should not be recommended.


The NASCIS III was a double-blind, randomized, controlled trial involving 499 patients with acute spinal cord injury. At 6-week and 6-month follow-up, a significant neurologic recovery persisted with the 48-hour methylprednisolone regimen that had been started within 3–8 hours after injury compared with the 24-hour regimen (neurologic change scores at 6 weeks, 7.6 vs 12.5, p=0.04; at 6 months, 11.2 vs 17.6, p=0.01). No difference in recovery was found when treatment was started within 3 hours of injury. In a third study group, tirilazad demonstrated efficacy similar to that of the 24-hour methylprednisolone regimen and was well tolerated. The authors concluded that methylprednisolone therapy should be continued for 24 hours if started within 3 hours of injury, but continued for 48 hours if started 3–8 hours after injury. (Class I)


In the NASCIS I trial, methylprednisolone in dosages up to 1000 mg/day did not improve neurologic recovery (sensory or motor function) in 330 patients with acute spinal cord injury. This article describes NASCIS II, a double-blind, placebo-controlled trial involving 487 patients treated within 8 hours of injury. Treatment consisted of a high-dose methylprednisolone intravenous bolus of 30 mg/kg followed by continuous infusion of 5.4 mg/kg/hour for 23 hours. Significant motor function and sensation were maintained at 6 weeks and 6 months with methylprednisolone compared with placebo or a naloxone bolus of 5.4 mg/kg followed by infusion of 4 mg/kg/hour for 23 hours. Complication rates were similar among the three groups. (Class I)

Head Injury

CRASH Trial Collaborators. Effect of intravenous corticosteroids on death within 14 days of 10008

The Medical Research Council’s Corticosteroid Randomization After Significant Head Injury (MRC CRASH) trial was a multicenter, randomized, placebo-controlled trial involving 10,008 adults with severe head injury. The goal was to determine the effect of early administration of methylprednisolone on risk of death at 2 weeks or disability at 6 months. Patients were randomized to receive intravenous methylprednisolone 2 g followed by 0.4 g/hour for 48 hours, or placebo, within 8 hours of injury. The study was stopped prematurely because an interim analysis showed a significant increase in relative risk of all-cause death in the methylprednisolone-treated group (RR 1.18, 95% CI 1.09–1.27, \( p=0.0001 \)). The relative increase in mortality in the methylprednisolone group did not differ by the severity or time of injury. The exact mechanism for the increased mortality is unknown. This is the first study that clearly refutes the mortality benefit of corticosteroids in the treatment of patients with severe head injury. (Class I)


Mannitol is an osmotic diuretic used to control raised intracranial pressure in patients with severe head injury. This is a task force–based guideline for treatment of traumatic intracranial hypertension with mannitol. The authors suggest that a bolus dose may be more effective than a continuous infusion. In addition, they advocate keeping serum osmolality below 320 mOsm/kg and avoiding hypovolemia to minimize the risk of acute tubular necrosis.


In patients with severe head injury, uncontrolled intracranial hypertension is associated with a high risk of mortality (84–100%). This article consists of guidelines for the use of barbiturates in treating such patients. In hemodynamically stable patients, high-dose barbiturates may be administered to lower intracranial pressure refractory to conventional medical or surgical treatment. Efficacy has not been proven for prophylactic lowering of intracranial pressure. These guidelines were based on a task force review.


Steroids have been investigated as a means to control raised intracranial pressure in patients with head trauma. This task force review did not support steroid administration because most of the available evidence does not suggest improved outcomes or lowered intracranial pressure.


These guidelines were based on a task force review of the literature on anticonvulsants for posttraumatic seizure prophylaxis. The authors concluded that phenytoin and carbamazepine decreased the risk of early posttraumatic seizure (defined as seizure onset within 7 days of injury). Improved survival benefit was not proven. The authors suggested valproic acid as an alternative agent, although it has been associated with a slightly higher mortality.


According to a secondary analysis of a prospective, randomized, double-blind, placebo-controlled trial, improved survival was not associated with the reduction of early posttraumatic seizures (within 7 days of injury) in 404 patients receiving phenytoin. Hypersensitivity reactions to phenytoin, primarily an idiosyncratic maculopapular rash, increased from 0.6% during week 1 to 2.5% during week 2. Overall, short-term phenytoin therapy was well tolerated. The authors suggest that mortality in patients who developed posttraumatic seizures might be linked to increased severity of the original traumatic brain injury. (Class II)


Seizures are common in patients with recent traumatic brain injury. This randomized, controlled trial compared a 7-day course of phenytoin (132 patients) with a 1-month (120 patients) and 6-month (127 patients) course of valproate for prevention of seizures associated with head trauma. Differences in the frequencies of seizures, mortality, and adverse effects were not statistically significant between groups. A trend toward higher mortality was found in the valproate group; however, none of the deaths...
were attributable to the most common adverse effects reported—liver dysfunction and thrombocytopenia. The authors concluded that valproate should not be routinely used for prophylaxis of posttraumatic seizures. (Class I)


This randomized, double-blind, placebo-controlled study is the landmark article supporting short-course phenytoin therapy for seizure prophylaxis in patients with severe head injury. Phenytoin (20-mg/kg load) or placebo was administered within 24 hours of injury, and maintenance doses were continued for 12 months. Phenytoin doses were adjusted to maintain free phenytoin levels within a target range of 0.75–1.5 mg/L. Patients were followed for 24 months. One week after injury, the seizure rate was 3.6% and 14.2% in the phenytoin and placebo groups, respectively (p<0.001). The differences in the frequency of late-onset seizures (day 8–year 2) between groups was not statistically significant (p>0.2). The authors concluded that phenytoin is effective only for prevention of early-onset (< 7 days) seizures after severe head injury. (Class I)

**Hemorrhagic Stroke**


Factor VII is beginning to play a role in the area of neurocritical care. A review of the pharmacology, pharmacokinetics, and treatment strategies in the critically ill neurosurgical patient population is discussed. The authors also review recently published case studies. This article provides a nice overview of factor VII treatment options in neurocritical care.


Vasospasm, a major complication after subarachnoid hemorrhage, is associated with high morbidity and mortality. Hypervolemia, hypertension, and hemodilution (triple-H) therapy is accepted as a standard of care; however, no randomized, controlled trials have been conducted to prove efficacy in the treatment of vasospasm. The authors review the literature and discuss the rationale for triple-H therapy.


A task force was formed to develop practice guidelines for the management of intracerebral hemorrhage. These guidelines provide recommendations for diagnosis, acute treatment, medical management, surgical treatment, and prevention of intracerebral hemorrhage. Algorithms for the management of blood pressure and intracranial pressure, parameters for fluid management, and recommendations for seizure prophylaxis and temperature regulation are provided.

**Ischemic Stroke**


These guidelines, which update the original 1994 guidelines, are intended to guide clinicians in the diagnosis and management of patients during the first 24–48 hours after ischemic stroke. Recommendations are provided for diagnostic testing and examination, general supportive care and treatment of complications, use of thrombolysis, role of anticoagulation, surgical interventions, and management of neurologic complications. Each recommendation is graded based on the level of supporting evidence.


The aim of this trial was to report the clinical outcomes and adverse effects of tissue-type plasminogen activator when used in practice. The 30-day clinical outcomes of intravenous tissue–type plasminogen activator 0.9 mg/kg, maximum 90 mg, were investigated in 389 consecutive patients with acute ischemic stroke. The 30-day mortality was 13%; in addition, 35% of patients had a very favorable outcome and
43% were functionally independent (modified Rankin scores of ≤ 1 and ≤ 2, respectively). Safety data from this observational study were consistent with the low frequency of symptomatic intracranial hemorrhage found in the National Institute of Neurological Disorders and Stroke trial. Caution should still be exercised for patients with major early infarction. (Class II)


Recombinant tissue–type plasminogen activator originally was studied when administered within 3 hours of symptom onset in patients with acute ischemic stroke. However, only a few patients received the drug within this time frame. In this randomized, placebo-controlled trial, alteplase was initiated in 3–5 hours after symptom onset. Outcomes were neurologic recovery at 90 days, functional outcome measures at 30 and 90 days, and serious adverse events. Neurologic recovery, functional recovery, and mortality were similar in both groups. Asymptomatic, symptomatic, and fatal intracranial hemorrhages were significantly higher in the alteplase group (p<0.05). The authors concluded that there was no benefit to administering alteplase 3–5 hours after symptom onset. (Class I)


The primary objective of this multicenter trial was to determine whether alteplase 0.9 mg/kg given within 6 hours of symptom onset would improve overall function in 800 patients with acute ischemic stroke. Efficacy of alteplase was independent of administration time (no difference when given < 3 or 3–6 hrs after stroke). No significant difference in efficacy or mortality was observed between alteplase and placebo at the 30- and 90-day end points. As seen in the first European Cooperative Acute Stroke Study (ECASS I) and the National Institute of Neurological Disorders and Stroke (NINDS) trial, symptomatic intracranial hemorrhage was higher in the alteplase group (8.8%) than in the placebo group (3.4%). (Class I)


Administration of tissue plasminogen activator (t-PA) has been associated with increased risk of intracranial hemorrhage when used to treat ischemic stroke. The authors investigated whether any variables collected in the NINDS t-PA stroke trial were associated with intracranial hemorrhage. Severity of neurologic deficit (OR 1.8, 95% CI 1.2–2.9) and brain edema and mass effect by computed tomography (OR 7.8, 95% CI 2.2–27.1) were the only independent variables associated with intracranial hemorrhage. Patients with severe neurologic deficits had more favorable outcomes at 90 days in the treatment group than in the placebo group (OR 4.3, 95% CI 1.6–11.9). The authors concluded that these patients were still candidates for t-PA if the agent was administered within 3 hours of symptom onset. (Class II)


Previous trials have suggested that early intervention with thrombolytic therapy improves clinical outcomes in patients with acute ischemic stroke, although the therapy is associated with a risk of symptomatic intracranial hemorrhage. In this prospective, randomized, double-blind, placebo-controlled trial, 624 patients were treated with recombinant tissue plasminogen activator (rt-PA) 0.9 mg/kg, maximum 90 mg. The results demonstrated significantly decreased neurologic deficit at 3 and 12 months in patients treated within 3 hours of stroke onset (OR 1.7, 95% CI 2–2.6). Symptomatic intracranial hemorrhage occurred more frequently in the rt-PA group (6.4%) than in the placebo group (0.6%) during the first 36 hours (p<0.001). No difference in mortality was detected. (Class I)


Restoration of cerebral blood flow is the goal of thrombolytic therapy in stroke patients. This randomized, controlled trial compared rt-PA with placebo in 620 patients with acute ischemic stroke. The patients presented within 6 hours of symptom onset and had experienced a stable
patients receiving pentobarbital, propofol, or midazolam were included in the analysis. Of the 193 patients, 48% died, with no differences based on the treatment received. Although pentobarbital was usually titrated to electroencephalogram background suppression, propofol and midazolam were titrated to the dosage needed to achieve seizure suppression. Overall, the limited data reviewed in this article suggest that treatment with pentobarbital or titration of any agent to electroencephalogram background suppression may be the most effective strategy for managing refractory status epilepticus. However, there was no obvious improvement in mortality. Prospective, controlled evaluations of this issue are needed.


This article provides a concise and clinically useful review of the clinical presentation, pathophysiology, initial management, and pharmacologic treatment of status epilepticus. The review contains a treatment algorithm that is widely employed in the management of patients with status epilepticus.


In this multicenter, randomized, double-blind trial, four intravenous regimens were compared for in-hospital treatment of generalized convulsive status epilepticus. A total of 570 patients were randomized to receive diazepam 0.15 mg/kg followed by phenytoin 18 mg/kg, lorazepam 0.1 mg/kg, phenobarbital 15 mg/kg, or phenytoin 18 mg/kg. Treatment success was defined as the absence of motor and electrical activity within 20 minutes of drug infusion, with no evidence of recurrence over the next 40 minutes. Intent-to-treat analysis revealed no statistically significant differences among regimens. For 384 patients with overt generalized convulsive seizures, lorazepam was more effective than phenytoin \((p=0.02)\) but did not differ from the diazepam-phenytoin and phenobarbital regimens. Based on at least comparable safety and efficacy for the four regimens, the authors concluded that for patients with generalized convulsive status epilepticus, lorazepam should be the initial treatment due to its ease of administration and improved response compared with phenytoin. (Class I)
Nutrition


This guideline was developed by a work group of EAST. Each of the six guideline subsections (route, timing, site, macronutrients, monitoring, and type of nutritional support) is a freestanding, evidence-based document. References and evidence tables associated with the guideline can be obtained on the EAST Web site (http://www.east.org). The guideline contains an algorithm summarizing the nutritional management of trauma patients.


The primary end point of this before-and-after, prospective investigation (100 patients in each phase) was to evaluate the impact of a nutritional management protocol on the time to initiation of feeding of patients in two medical-surgical ICUs. The protocol was evidence based but lacked high-level evidence for some recommendations. Enteral versus parenteral nutrition, ability to reach caloric goals, and length of ICU stay were evaluated. Patients in the postimplementation group were more likely than those in the preimplementation group to be fed enterally, and their mean duration of mechanical ventilation was decreased. However, the results were statistically significant (p=0.009 and p=0.03, respectively) only after adjusting for a variety of covariates, such as baseline nutritional status and severity of illness. The logic behind this adjustment was questioned in an editorial accompanying the article (Zaloga GP, Bortenschlager L. Anorexia protocolis. Chest 2004;125:1195–7). No other significant differences between groups were noted, probably because of the comparable number of patients receiving enteral feeds in the preprotocol and postprotocol groups (68% and 78%, respectively, p=0.08). This study is notable because of its prospective design and comprehensive approach to the nutritional management of critically ill patients. (Class II)


Based on an extensive review of the literature, an interdisciplinary, technical advisory group of nutrition support specialists updated the 1993 guidelines to assist clinicians in managing patients in both outpatient and inpatient settings. Each topic is presented in the same format: background, evidence, special considerations (if applicable), and graded recommendations for specialized nutrition support.


The appropriate indications for the use of enteral feeding formulas supplemented with immune-altering nutrients in the ICU are controversial despite the availability of consensus guidelines (Kudsk KA, Moore F, Martindale RG, Cresci G, McClave S, Schloerb PR. Consensus recommendations from the U.S. summit on immune-enhancing enteral therapy. J Parenter Enteral Nutr 2001;25(suppl):S61–2). This meta-analysis is important not only because it evaluates the use of such formulas in critically ill patients, but also because it raises important questions and concerns regarding research in this area.


This meta-analysis of 26 randomized trials evaluated morbidity and mortality outcomes in 2211 surgical or critically ill patients receiving total parenteral nutrition (TPN). Although TPN did not improve mortality, a trend toward decreased complication rates was found in patients classified as malnourished.


Outcome data for morbidity and mortality in surgical patients requiring TPN are lacking in the published literature. The primary objective of this prospective, randomized, controlled trial was to determine whether perioperative TPN decreased serious complications secondary to major abdominal or thoracic surgery. A total of 395 patients were randomized—192 to the TPN group and 203 to the control group. Rates of major complications, mortality, and noninfectious complications were not statistically significant between the two groups. Patients receiving TPN had more infectious complications
than controls (14.1% vs 6.4%, RR 2.20, 95% CI 1.19–4.05). Severely malnourished patients, however, experienced an overall benefit from TPN since a significantly lower rate of infectious complications was found in this group (RR 0.12, 95% CI 0.02–0.91). (Class I)

Sedation, Analgesia, Delirium, and Neuromuscular Blockade


These authors tested the recommendation from the guidelines of the Society of Critical Care Medicine and the American Society of Health-System Pharmacists for routine use of a peripheral nerve stimulator with continuous neuromuscular blockade. Medical intensive care patients were randomized to train-of-four every 4 hours versus clinical assessment during cisatracurium infusions of 50–70 hours. Cisatracurium was titrated to maintain one or two of four twitches. No difference was demonstrated in time to recovery of neuromuscular function (four of four twitches) or dosing requirement for the 30 patients studied. The authors suggested that routine use of train-of-four was not needed during cisatracurium infusion. The article and its accompanying editorial (Sessler CN. Train-of-four to monitor neuromuscular blockade? Chest 2004;126:1018–22) correctly suggest that these data cannot be extrapolated to infusions of aminosteroidal neuromuscular blockade agents, whereas accumulation of active metabolites may contribute to prolonged drug effects. (Class I)


The authors developed a hypothesis for the development of a syndrome characterized by cardiac failure, rhabdomyolysis, severe metabolic acidosis, and renal failure after prolonged or high-dose propofol infusion. This syndrome appears to be rare; only 21 pediatric and 14 adult cases have been reported in the literature. Most of the reports involve patients with neurologic illnesses or acute inflammatory diseases (e.g., severe sepsis) as priming factors in combination with triggering factors, such as concurrent catecholamine infusions or steroids. Propofol doses greater than 5 mg/kg/hour for more than 48 hours were reported in the cases described.


The RASS is a modification of the Riker Sedation-Agitation Scale, expanded to score 10 levels rather than the original seven. In particular, the degrees of light sedation are expanded to differentiate response to voice from response to physical stimulation. This prospective cohort trial evaluated the RASS used in adult patients in medical and coronary ICUs compared with the Glasgow Coma Scale and Ramsay scale. The study serves as an independent validation of the utility of the RASS. In 290 paired observations by nurses, the RASS and Ramsay scale both demonstrated excellent interrater reliability as well as criterion, construct, and face validity. Both were superior to the Glasgow Coma Scale. The RASS detected changes in sedation status over consecutive days in patients with varying levels of consciousness and delirium. Scores changed appropriately with dosages of sedatives and analgesics. (Class II)


This is the second observational pain scale to be published and validated at the site of origin. The nonverbal pain scale yields a score of 0–2 for each of the major categories (facial expression, activity [movement], guarding, vital signs, and skin appearance), for a potential score of 0–10. Validation occurred with 100 paired assessments on all three 8-hour nursing shifts, although most tests were performed during the day. The nonverbal pain scale was compared with a pediatric scale for face, legs, activity, cry, and consolability due to the lack of a previous gold standard for assessment of nonverbal patients. Interrater reliability for the nonverbal pain scale was good (α=0.78). However, no assessment was performed to ensure that the change on the nonverbal pain scale was correlated with different degrees of pain. (Class II)

This introductory document accompanies two corresponding guideline documents (described below) developed jointly by the American Society of Health-System Pharmacists and the Society of Critical Care Medicine. Levels of evidence and definitions for the grades of recommendation are described in this document. The three articles in this series have also been published elsewhere (Am J Health-Syst Pharm 2002;59:147–95).


This guideline, second in the series of three articles, summarizes the extensive literature on analgesia and sedation of critically ill patients with emphasis on patient assessment, therapeutic options, and recommendations for optimal therapy. Examples of assessment tools are provided along with an algorithm approach to treatment plan development. Associated topics include management of delirium and issues related to nonpharmacologic therapy and sleep.


The third article in this series similarly summarizes the literature on the use of neuromuscular blocking agents in critical care patients. The authors describe the mechanism of action and the pharmacology of the individual agents. They also describe the appropriate indications and recommended agents, along with suggestions for how to monitor for effectiveness and adverse effects. The article also reviews the syndromes and risk factors leading to delayed muscle recovery after treatment with these agents. A template for evaluation of the pharmacoeconomic impact of individual agents is provided.


In this double-blind clinical trial, 24 mechanically ventilated patients were randomized after surgery to receive sedation with lorazepam or midazolam by continuous infusion for 12–72 hours. Sedation was titrated through infusion rate changes with a goal of Ramsay scale level 3–4. Serum concentrations and pharmacodynamic effects were also measured. The estimated sedative potency of lorazepam was twice that of midazolam on a milligram-to-milligram basis. The relative amnesic potency of lorazepam was 4 compared with midazolam. Lorazepam was associated with longer times to emergence from 72 hours of sedation with both deep (31.1 hrs) and light (11.9 hrs) regimens compared with deep and light regimens of midazolam (14.9 and 3.9 hrs, respectively). However, excluding patients with renal or hepatic insufficiency may have minimized accumulation of midazolam’s active metabolites that could alter awakening times. (Class I)


Intensive care patients can experience prolonged weakness after a critical illness such as sepsis. Myopathy treated with corticosteroids has been well described, and the potential contribution of neuromuscular blocking agents was well documented in the neuromuscular blocking agent guidelines. In this article, the authors studied 73 patients with sepsis and assessed neuromuscular changes with electrophysiologic studies on the days 10 and 21 of mechanical ventilation. Multivariate logistic regression analysis of the risk factors for critical-illness polyneuropathy demonstrated that hyperosmolality, parenteral nutrition, neuromuscular blocking agents, and neurologic failure were independently associated with critical-illness polyneuropathy, whereas renal replacement therapy appeared protective. Forty six patients had critical-illness polyneuropathy on the first assessment, with an additional four diagnosed at the second evaluation. As demonstrated by others, critical-illness polyneuropathy contributed to a longer duration of mechanical ventilation and hospital stay. (Class II)


In this study, 128 critically ill, mechanically ventilated patients were randomized to daily interruption of sedative infusion or routine sedative titration. Patients were further randomized to therapy with propofol or midazolam.
Duration of mechanical ventilation was reduced from 7.3 to 4.9 days (p=0.004) with daily sedative interruption. Length of stay in the ICU was also significantly reduced. Additional information was reported in subsequent publications (Kress JP, Pohlman AS, Hall JB. Effects of sedative interruption in critically ill, mechanically ventilated patients receiving midazolam or propofol. J Clin Outcomes Manage 2001;8:33–9; and Kress JP, Gelbach B, Lacy M, et al. The long-term psychological effects of daily sedative interruption on critically ill patients. Am J Resp Crit Care Med 2003;168:1457–61). (Class I)

Toxicology


This two-part series provides a comprehensive review of the evaluation, treatment, and outcome of common overdoses and intoxications from the perspective of the critical care practitioner. The first review focuses on the general management of the intoxicated patient, including physical examination and laboratory findings of common toxidromes, nonspecific supportive therapies, and detoxification. The second review discusses the management of specific intoxicants and poisons.


This article provides a concise and well-written review of current strategies for management of common overdoses and poisonings in the critical care setting. Gastrointestinal decontamination and methods for enhancing elimination of the toxin are covered, and brief overviews are provided for the most common poisonings encountered in the ICU.

Vasoactive Agents


This recent article is an evidence-based review by the American College of Critical Care Medicine of the literature related to hemodynamic support in patients with sepsis. Specific recommendations are provided for monitoring parameters and end points for therapy. Areas discussed include fluid resuscitation, vasopressor therapy, and inotropic therapy for adult patients with sepsis.


This concise, well-written review provides an overview of the management of septic shock. Principles of fluid resuscitation, vasopressor and inotropic drug treatment, and the role of vasopressin are covered. A one-page flow diagram of management decisions based on hemodynamic and clinical response is also provided.


The increased use of vasopressin in the management of vasodilatory shock is based primarily on small case series, case reports, and very small prospective studies. This study is the largest prospective, randomized, controlled study of vasopressin published to date. Forty-eight patients with vasodilatory shock were randomized to receive arginine vasopressin at 4 U/hour (0.067 U/min) in addition to norepinephrine or continued norepinephrine therapy. The vasodilatory shock in all patients was due to postcardiotomy syndrome (41.6%), systemic inflammatory response syndrome (29.2%), or sepsis (29.2%). In addition, patients remained hypotensive (mean arterial pressure < 70 mm Hg) despite adequate fluid resuscitation and required norepinephrine administered at more than 0.5 µg/kg/minute. In the arginine vasopressin group, heart rate decreased significantly; mean arterial pressure increased and remained significantly higher; and cardiac index, stroke volume index, and left ventricular stroke work index were increased compared with the norepinephrine group. Norepinephrine requirements were significantly lower in the arginine vasopressin group. New-onset tachycardic atrial fibrillation occurred in two (8.3%) patients in the arginine vasopressin group, compared with 14 (54.3%) patients in the norepinephrine group (p<0.001). Regional partial pressure of carbon dioxide (PCO₂) tension and PCO₂ gap, as measured by gastric tonometry, were significantly
lower in the arginine vasopressin group. This suggests improved gastrointestinal perfusion in the intervention group. Mortality in the ICU was 70% in both study groups. Overall, arginine vasopressin was safe and effective for management of norepinephrine-dependent vasodilatory shock and led to reduced norepinephrine dosage requirements, improved global and regional hemodynamics, and fewer tachyarrhythmias. (Class I)


The authors reviewed the 30-year history of low-dose dopamine treatment in the ICU. Based on the collective evidence, they concluded that this therapy does not have a beneficial effect in critically ill patients with oliguria. In addition, substantial evidence indicates that low-dose dopamine may have significant detrimental effects. The authors concluded that there is no justification for the use of low-dose dopamine in treating critically ill patients.


This prospective, cohort, observational study was designed to identify factors associated with outcome in 97 patients treated for septic shock. Nineteen clinical, biologic, and hemodynamic variables were collected during the first 48–72 hours of treatment. Stepwise logistic regression analysis was used to identify variables that were independently and significantly associated with outcome. The primary outcome variable was hospital mortality, which was 73% in this cohort. Five factors were significantly associated with outcome. Lactic acidosis, multiple organ failure, pneumonia, and oliguria were associated with an increased risk of death, and norepinephrine as a component of hemodynamic management was associated with reduced risk of death (relative risk ratio 0.68, 95% CI 0.54–0.87) when compared with high-dose dopamine or epinephrine. Currently, no mortality outcome data from prospective randomized trials evaluate the hemodynamic management of septic shock. This observational study is the only study reporting that mortality outcome may be affected by the choice of vasopressors. The study was limited by its observational design and relatively small sample. (Class II)

Economic Benefits of Intensive Care Unit Pharmacy Services


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