Polyclonal intravenous immunoglobulin for the prophylaxis and treatment of infection in critically ill adults

Kevin B Laupland MD FRCPC

Infection is a major cause of morbidity and mortality in critically ill patients. Despite advances in technology, its mortality rate has changed minimally over the past two decades, and new therapies are needed. Polyclonal intravenous immunoglobulin (IVIG) has been investigated both as a preventive and a treatment modality for sepsis and septic shock in critically ill adult patients. Prophylaxis with IVIG has been shown to reduce significantly the incidence of infection, particularly pneumonia, in selected postsurgical intensive care patients. However, it does not reduce mortality. The risk-benefit and cost effectiveness of this therapeutic intervention have not been determined, and its routine use is therefore not recommended. Treatment with IVIG has been shown in a number of small trials and a meta-analysis to reduce dramatically sepsis and septic shock mortality. However, a large, unpublished randomized trial has apparently shown no mortality benefit with this therapy. Despite limited evidence, IVIG has become the standard of care for the management of group A streptococcal toxic shock syndrome. At present, clinical equipoise exists for the use of IVIG in the treatment of sepsis and septic shock, and further study is needed.

Key Words: Critical care; Infection; Intensive care unit; Intravenous gamma globulin; Sepsis; Septic shock

Infection is a major cause of morbidity and mortality in critically ill patients. Although Canadian incidence rates are unknown, it is estimated that 300,000 to 500,000 septic episodes occur each year in the United States, with a crude mortality rate of 35% (1). Despite remarkable advances in medical and surgical care, the mortality rate from severe infections has declined only minimally over the past few decades (2). As a result, considerable effort has been devoted to developing new sepsis and septic shock therapies.

The dramatic clinical manifestations of sepsis and septic shock are believed to arise from an exaggerated immune response to Gram-negative lipopolysaccharide (LPS), or Gram-positive lipoteichoic acid or protein exotoxins. These bacterial antigens and superantigens induce the host production of proinflammatory cytokines, most importantly, interleukin-1-beta (IL-1β) and tumour necrosis factor-alpha (TNF-α), which then lead directly and indirectly to the clinical manifestations of sepsis and septic shock (3,4). New therapies have therefore targeted both bacterial components and the host-derived immune mediators in the management of sepsis and septic shock.

Large randomized clinical studies investigating the use of IL-1β receptor antagonists, soluble TNF-α receptors, and monoclonal anti-LPS and anti-IL-1β antibodies have not demonstrated any significant mortality benefit (5,6). With the exception of one recent trial that enrolled highly selected patients, studies using anti-TNF-α antibody have also shown no mortality benefit (7). Alejadria et al (8) performed a meta-analysis of intravenous antibody therapies in sepsis and septic shock, and found no overall mortality benefit from monoclonal antiantidotoxin (n=1736) or anticytokine (n=4318) products. There are a number of possible explanations for their apparent lack of efficacy. The timing of administration and dosing of these agents may not have been optimal. In addition, it may have been necessary to treat patients with therapies directed at both the inciting antigens and a number of different cytokines simultaneously rather than using monotherapy. This may be because the pathogenesis of sepsis and septic shock involves a complex interaction between the organism and a broad range of host mediators. Another possibility is that the agents studied were directed at the wrong targets or failed to address the importance of the interactions between the immune and coagulation systems in sepsis. The recent clinical trial showing that recombinant human activated protein C reduced mortality in patients with severe sepsis supports this argument (9). Because it has a broad range of known and unknown specificities and activities, intravenous immunoglobulin (IVIG) has been investigated as a preventive and therapeutic modality for infections in the critically ill. The present review article reviews the rationale and evidence for the use of IVIG in the prophylaxis and treatment of infection in critically ill adult patients.

**WHAT IS IVIG?**

IVIG is polyclonal immunoglobulin that is derived from large pools of normal donor serum and prepared for intravenous injection. Intramuscular preparations of immune globulin first became available in the 1950s; IVIG became available in 1979 (10). Standard IVIG contains primarily immunoglobulin G from as many as 30,000 plasma donors (11,12). As a result of pooling, IVIG reflects the prevalent immunity of the plasma donor population and, therefore, antibody titres and specificities vary among different lots and manufacturers (13). Different preparations have in common a broad range of antibodies to microorganisms and their extracellular products, cytokines, and contain soluble human leukocyte antigens class I and II (14).

Therapeutic IVIG is generally well tolerated and has a good safety profile. A 2 g/kg dose will increase serum levels by 2 to 3 g/L (normal range 6 to 16 g/L), with a return to baseline at 23 to 28 days (10). There are anecdotal reports of acute tubular necrosis, aseptic meningitis, central retinal vein obstruction, myocardial infarction and deep venous thrombosis associated with IVIG use. However, large randomized trials investigating IVIG use in patients with a wide range of medical conditions have shown a low incidence of adverse events that is comparable with use of a placebo (12,15-17). Because it is a concentrated colloid solution, high doses of IVIG may precipitate congestive heart failure or exacerbate volume overloaded states. Patients with selective immunoglobulin A deficiency may have a severe anaphylactic reaction to IVIG, and is contraindicated in these patients. Compared with other blood and plasma products, the risk of transmitting infectious agents with IVIG is believed to be negligible. This is because, in addition to careful screening of donors and rigorous quality control, the purification process is toxic to viruses and bacteria. The process varies among manufacturers, but is usually based on the cold ethanol fractionation method of Cohn and Onley (13,18). Additional processing is then performed to remove aggregates and anticomplement proteins. Techniques may include enzymatic cleavage, chemical modification and nonmodified purification procedures, including polyethylene glycol/hydroxethyl starch precipitation, lyophilization, dialfiltration, ultrafiltration, ion exchange, and exposure to acid, heat or detergents (13,18). Despite these precautions, there is always the potential risk of transmitting infectious agents that have yet to be discovered.

IVIG has been used therapeutically in a broad range of infectious and noninfectious disorders. However, it is expensive, with an approximate cost of $40/g, the evidence for its use in many indications is weak, and there is an ever present risk of shortages because Canada imports more than one-half of its IVIG supply (19). Because of these concerns, a Canadian consensus group was formed to explore indications for IVIG use. They conducted a survey among Canadian specialists who reported a total of 142 potential uses for IVIG (10). The investigators then performed an extensive review of the literature to determine evidence-based indications for IVIG use. However, with the exception of streptococcal toxic shock syndrome, they did not present the evidence for IVIG as a preventive or treatment modality for sepsis and septic shock in adults.
TABLE 1
Randomized, controlled trials of intravenous immunoglobulin (IVIG) for the prophylaxis of infection in critically ill adults

<table>
<thead>
<tr>
<th>Setting</th>
<th>Design (doses used)</th>
<th>Patients (n)</th>
<th>Main outcomes</th>
<th>Limitations</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surgical ICUs</td>
<td>RDBPCT (standard IVIG 400 mg/kg, LPS IVIG 400 mg/kg or 25% albumin)</td>
<td>329</td>
<td>Infection incidence with standard IVIG 33% versus placebo 47%, P=0.03; fewer pneumonias, shorter ICU stay with IVIG compared with placebo</td>
<td>Analysis, not intention-to-treat, but unlikely to change results</td>
<td>11</td>
</tr>
<tr>
<td>Trauma patients in ICU</td>
<td>RDBPCT (36 g IVIG versus 0.03% albumin)</td>
<td>150</td>
<td>Decreased overall incidence of infection 47% versus 68%, P=0.02; decreased antibiotics in IVIG group</td>
<td>Blinding process not well described</td>
<td>12</td>
</tr>
<tr>
<td>Postoperative open heart surgery patients with cutaneous anergy preoperatively</td>
<td>RDBPCT (20 g IgA- and IgM-enriched IVIG versus saline)</td>
<td>40</td>
<td>Infection incidence 5% IVIG versus 43% placebo, P=0.007</td>
<td>Highly selected patients (40 of 515)</td>
<td>20</td>
</tr>
<tr>
<td>Trauma patients in ICU with injury severity score 16 to 50</td>
<td>RDBPCT (1 g/kg IVIG versus 1 g/kg albumin)</td>
<td>39</td>
<td>No difference in overall infection rates, but fewer pneumonias and noncatheter-related infections; no difference in ICU length of stay or antibiotic use</td>
<td>Small numbers may not detect significant differences; highly selected patients (39 of 196)</td>
<td>21</td>
</tr>
<tr>
<td>Major surgery and trauma ICU patients</td>
<td>RDBPCT (40 g IVIG versus 5% dextrose)</td>
<td>40</td>
<td>No difference in rates of sepsis or mortality; less positive blood cultures in IVIG group in the second week of observation</td>
<td>Small numbers may not detect significant differences; incidence of nonseptic infections not reported</td>
<td>23</td>
</tr>
</tbody>
</table>

ICU Intensive care unit; Ig Immunoglobulin; LPS Lipopolysaccharide; RDBPCT Randomized, double-blinded, placebo controlled trial

EVIDENCE FOR IVIG AS PROPHYLACTIC THERAPY

Randomized clinical trials investigating the use of IVIG as a prophylactic agent in critically ill patients were initially identified by searching MEDLINE from 1966 to December 2000. Search terms were immunoglobulins, intravenous immunoglobulins, passive immunization, or immunoglobulin G and infection, sepsis, sepsis syndrome or septic shock. Studies included were limited to those that enrolled patients older than 12 years, and human and clinical trials. Abstracts were then screened to identify prospective clinical trials in critically ill patients that compared an IVIG therapy group with a control group. Full-length reports were then retrieved, and further studies were searched for using the bibliographies of these reports and review articles. Five prospective clinical trials that focused on the use of IVIG as a preventive therapy for infection in intensive care unit (ICU) patients were identified for critical appraisal (Table 1).

The best quality evidence for the efficacy of IVIG in the prophylaxis of infection has been reported by The Intravenous Immunoglobulin Collaborative Study Group (11). This randomized, double-blind, placebo controlled trial conducted in surgical ICUs in Belgium, Switzerland and the United States randomly assigned 352 patients (329 evaluated) to treatment with standard IVIG, LPS hyperimmune IVIG or 25% albumin placebo. They observed a lower rate of overall infection in the standard IVIG group of 36 of 109 patients (33%) compared with 53 of 112 placebo patients (47%, P=0.03). Patients treated with standard IVIG compared with placebo also had lower rates of pneumonia (14% versus 27%, respectively, P=0.04) and a shorter median length of stay in the ICU (four versus six days, respectively, P=0.02). There was no significant difference among the groups with respect to the incidence of septic shock or mortality rate. Hyperimmune LPS IVIG was similar to placebo for all comparisons.

Glinz et al (12) from Zurich, Switzerland reported a randomized, double-blind, placebo controlled trial of 150 trauma patients mechanically ventilated for longer than 24 h. Seventy-six patients were allocated to 36 g of IVIG, and 74 patients were allocated to 0.03% albumin. The number of infections (over 42 days) was 36 in the IVIG group (47%) versus 50 in the control group (68%, P=0.02), with most of the difference secondary to a lower incidence of pneumonia in the IVIG group compared with the control group at 28 patients (37%) versus 43 patients (58%, P=0.001), respectively. No statistically significant differences were observed in the mortality rate of 23 patients (30%) versus 15 patients (20%), respectively, but IVIG use was associated with fewer antibiotic days compared with the control group (1120 versus 1641 days, respectively).

Kress et al (20) conducted a randomized, double-blind, placebo controlled trial in Wurzburg, Germany with anergic postoperative open heart surgery patients. They preoperatively tested the cutaneous delayed-type hypersensitivity responses of 515 patients to seven common antigens and identified 44 patients with anergy. These patients were then randomly assigned to either 20 g IVIG (immunoglobulin A- and immunoglobulin M-enriched) or saline and assessed prospectively for the development of infection postoperatively. Of 40 evaluated patients, one patient (5%) in the IVIG group developed an infection...
compared with nine patients (43%) in the placebo group (P=0.007). No significant differences were observed with respect to mortality rate, or ICU or hospital length of stay, although the study was underpowered to determine clinically significant differences.

In a randomized, double-blind, placebo controlled trial by Douzinas et al (21) from Athens, Greece, 39 multitrauma patients were selected from 196 screened patients and were randomly assigned within 12 h of ICU admission to 1 g/kg of IVIG or human albumin divided over four days. Baseline characteristics were similar in the two groups, except that the IVIG group had higher injury severity scores (22). After controlling for injury severity scores, no overall difference in infection incidence was observed, but pneumonia (P=0.003) and noncatheter-related infections (P=0.04) were reduced with IVIG compared with the control group. Catheter-related infections, antibiotic use, mortality and ICU length of stay did not significantly differ between the groups. Limitations of this study included the small sample size and failure to report the incidence of nonseptic infectious complications.

Mao and colleagues (23) from Torino, Italy conducted a randomized, double-blind, placebo controlled trial among 40 surgical ICU patients. They randomly assigned postoperative major surgery and trauma patients to 40 g IVIG or 5% dextrose. Five IVIG-treated and three control patients died within 48 h of random assignment and were excluded from analysis. Three IVIG-treated patients developed sepsis compared with six control patients (P=0.05). Although there was no difference in the development of positive blood cultures during the first week of observation, in the second week, no patients treated with IVIG had positive blood cultures compared with six patients in the placebo group (P<0.05). The authors did not report the incidence of other infectious syndromes. There was a nonsignificant difference in mortality among patients treated with IVIG (two of 15, 13%) versus placebo (four of 17, 24%). This study was limited by its small sample size and failure to report the incidence of nonseptic infectious complications.

### Evidence for IVIG as Adjunctive Therapy

Polyclonal IVIG has been investigated as an adjunctive treatment modality in sepsis and septic shock. Randomized clinical trials of IVIG in the treatment of sepsis and septic shock were searched using the strategy described for prophylaxis in the preceding section. Eight articles were identified that fulfilled the criteria of prospective, controlled studies of IVIG treatment focusing on critically ill adults with sepsis and septic shock (Table 2).

### Table 2

<table>
<thead>
<tr>
<th>Setting</th>
<th>Design</th>
<th>Patients (n)</th>
<th>Main outcomes</th>
<th>Limitations</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surgical and trauma ICU patients with sepsis scores ≥20</td>
<td>RDBPCT (1 g/kg IVIG versus albumin)</td>
<td>62</td>
<td>Reduced mortality of 33% IVIG versus 67% placebo, P&lt;0.05</td>
<td>Highly selected surgical patients</td>
<td>24</td>
</tr>
<tr>
<td>Surgical and trauma ICU patients with sepsis scores ≥17</td>
<td>RDBPCT (1 g/kg IVIG versus albumin)</td>
<td>51 (excluding those pooled from prior study [24])</td>
<td>Reduced mortality of 29% IVIG versus 61% placebo, P=0.021</td>
<td>Highly selected surgical patients</td>
<td>27</td>
</tr>
<tr>
<td>ICU patients with Gram-negative septic shock</td>
<td>Open-label, randomized trial (60 g IVIG versus ‘usual care’)</td>
<td>55</td>
<td>Reduced septic mortality of 4% IVIG versus 32% control, P&lt;0.01</td>
<td>Highly selected patients (69 of 860), multiple interim analyses, unblinded</td>
<td>28</td>
</tr>
<tr>
<td>ICU patients with clinical sepsis and endotoxemia</td>
<td>Randomized, controlled trial (IVIG 0.5 g/kg versus control)</td>
<td>46</td>
<td>Nonsignificant decrease in mortality of 63% IVIG versus 86% control, P=not significant</td>
<td>Unclear if adequate blinding, intervention late in illness, small sample size, low dose IVIG</td>
<td>30</td>
</tr>
<tr>
<td>Postoperative patients with sepsis</td>
<td>Open-label, randomized trial (IVIG dose not stated)</td>
<td>35</td>
<td>Mortality rate 44% IVIG versus 76% control, P=not significant</td>
<td>Unblinded, small sample size</td>
<td>31</td>
</tr>
<tr>
<td>Septic medical and surgical ICU patients</td>
<td>Open-label, randomized trial (1 g/kg IVIG versus ‘usual care’)</td>
<td>24</td>
<td>No significant difference in mortality (58% IVIG versus 75% control)</td>
<td>Grossly underpowered, unblinded</td>
<td>32</td>
</tr>
<tr>
<td>ICU patients at first sign of infection</td>
<td>IVIG (dose not stated) versus ‘usual care’</td>
<td>104 (only 14 with sepsis or septic shock)</td>
<td>No difference in mortality</td>
<td>Unblinded, small number of patients with sepsis or septic shock</td>
<td>33</td>
</tr>
<tr>
<td>Septic medical and surgical ICU patients</td>
<td>Multicentred, RDBPCT (0.9 g/kg IVIG versus albumin)</td>
<td>653</td>
<td>No difference in mortality, but actual rates not reported</td>
<td>Unpublished despite completion four years ago, enrolment appears to have ended prematurely</td>
<td>34</td>
</tr>
</tbody>
</table>
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The best evidence for efficacy of IVIG in the treatment of sepsis and septic shock comes from an Italian trial by Dominioni et al (24). In this multicentred, randomized, double-blind, placebo controlled trial, they enrolled 62 consecutive surgical and trauma patients with sepsis scores higher than 20 (mean Apache II score 18) and randomly assigned them to either 1 g/kg of IVIG or albumin administered over a five-day period (25,26). They observed a 28-day mortality rate of 11 of 29 IVIG-treated patients (38%) compared with 22 of 33 control patients (67%), P<0.05. The decrease in mortality was related to a decrease in sepsis-related mortality in the IVIG group compared with the control group (7% versus 33%, respectively, P<0.05). A trend for a decrease in positive blood cultures after randomization was observed among IVIG patients, suggesting that this therapy may increase the clearance of microorganisms from the blood (five of 29 IVIG-treated patients versus 12 of 33 control patients, P>0.05).

Dominioni and colleagues (27) subsequently completed a second randomized trial similar to the first, with the exception that they included patients with sepsis scores of 17 or higher. They reported the results pooled with their initial trial and observed a lower overall mortality rate of 19 of 57 IVIG-treated patients (33%) compared with 36 of 56 control patients (64%, P<0.005) (24,27). The decrease in mortality was due primarily to a reduction in mortality among patients with intermediate sepsis scores of 20 to 25 (11 of 33 IVIG-treated patients [33%] versus 23 of 35 control patients [66%], P<0.025). No statistically significant differences were observed in mortality reductions with IVIG among patients with sepsis scores of 17 to 19 (none of 10 IVIG-treated patients versus two of seven control patients) and higher than 25 (eight of 14 IVIG-treated patients versus 11 of 14 control patients). An important finding of this trial was that a consistently lower mortality rate was observed with IVIG therapy in the second group of independently studied patients. If the patients in the first published trial by Dominioni and colleagues (24) are excluded from the analysis, the mortality rate among the 51 patients in the second study (27) is again statistically significant – eight of 28 IVIG-treated patients (29%) compared with 14 of 23 control patients (61%, P=0.021, Fisher’s exact test).

Schedel et al (28) reported an open-label, prospective, randomized, controlled trial of 55 patients with Gram-negative septic shock (medical and surgical) as identified by the limulus amebocyte lysate assay. They randomly assigned patients to 60 g IVIG or ‘usual care’. They observed a septic mortality rate of one of 27 IVIG-treated patients (4%) compared with nine of 28 control patients (32%) within six weeks of randomization (P<0.01). Furthermore, they observed a faster decrease in endotoxin in blood (24 h) in IVIG-treated patients compared with control patients (four days, P<0.01). This study was limited because patient allocation was not blinded, and because endotoxemia was an enrolment criterion, the results can only be generalized to patients with Gram-negative sepsis. Furthermore, this study has been criticized both in methodology and in conduct, because the authors performed multiple interim analyses, reported septic rather than all-cause mortality rate and may have modified the inclusion criteria after the start of the trial (29).

Grundmann and Hornung (30) from Köl, Germany reported on a randomized, controlled trial of IVIG (0.5 g/kg) therapy among 46 ICU patients with clinical sepsis and endotoxemia. They observed a nonsignificant lower mortality rate of 15 of 24 IVIG-treated patients (63%) compared with 19 of 22 control patients (86%). An important limitation is that they treated patients at least 48 h after presentation (average six days after ICU admission). This may potentially bias for no effect for IVIG, because if this treatment is useful, it is expected to have its greatest effect if given early in the disease course, when the cytokine cascade is initiated. The small study sample size and the use of a relatively low dose of IVIG may further bias the results toward no effect. Finally, the study methodology did not define whether there was adequate blinding, which could potentially bias the result in favour of or against IVIG efficacy.

This same group of investigators also reported, in the German literature, a single centre, open-label, randomized clinical trial (31). They randomly assigned 35 postoperative ICU patients with sepsis scores of 12 or higher to treatment with IVIG (total gram dose not specified) or control. There was no statistically significant reduction in mortality among patients treated with IVIG (eight of 18, 44%) compared with control (13 of 17, 76%). They also observed a nonstatistically different lower ICU stay (13 days versus 16 days) and ventilator days (10 days versus 13 days). The main limitations of this study relate to its small sample size and that the investigators were not blinded to the type of treatment received.

De Simone et al (32) reported a randomized, open-label clinical trial of IVIG 1.0 g/kg versus ‘usual care’ in 24 septic ICU patients (16 surgical) from Rome, Italy. They found a statistically nonsignificant mortality reduction in patients undergoing IVIG treatment compared with control patients (seven of 12 patients [58%] versus nine of 12 patients [75%], respectively). However, the study was grossly underpowered to detect a clinically significant mortality reduction. A trend was observed for an increased median survival of 30 days versus 10 days in the IVIG and control groups, respectively (P<0.1). Of interest, the percentage of days of ICU stay on antibiotics was less in the IVIG group compared with the control group (38% versus 95%, P<0.01). A conclusion regarding an antibiotic-sparing effect of IVIG cannot be made because the study was not blinded, and knowledge of treatment with IVIG may have influenced the decision to discontinue treatment with antibiotics.

Just and colleagues (33) reported, in the German literature, an open-label, randomized, controlled trial in 104 ICU patients. They randomly assigned patients at first sign of infection to treatment with IVIG (total gram dose not stated) or usual care. There was no significant difference in mortality among patients treated with IVIG (25 of 50,
50%) compared with control patients (22 of 54, 41%). No significant differences in overall length of stay or number of days of mechanical ventilation were observed. The most important limitation of this study is that patients tended to have mild infection-related illness, because only six patients treated with IVIG and eight control patients had criteria for either sepsis or septic shock (8). Thus, the severity of illness of the majority of the patients in this study was not related to infection, and IVIG would not be expected to modify significantly their disease course.

Overall, the published clinical trials suggest a beneficial mortality effect for treatment with IVIG in selected patients with sepsis and septic shock. Alejandra et al (8) have performed a meta-analysis based on five trials (24,28,30-32). These authors calculated an overall dramatic reduction in mortality among adult patients treated with IVIG compared with control patients (n=222, RR=0.60, 95% CI 0.47 to 0.77) (8). However, these results must be interpreted carefully, because each of the individual studies were relatively small, had highly selected patients, and were of variable quality in both methodology and conduct, as detailed in the preceding paragraphs.

Werdan and colleagues recognized the importance of a large, randomized trial to further define the role of IVIG in septic shock and completed a trial with 653 septic patients (34-36). In their multicentred, randomized, double-blind, placebo controlled trial, they enrolled medical and surgical patients with sepsis scores of 12 to 27 and Apache II scores of 20 to 35 (35). Patients were randomly assigned to 0.9 g/kg IVIG versus albumin placebo. This study was apparently completed in 1996, but it has not been published as a full manuscript (36). In a review article on immunoglobulin use in sepsis, the authors reported that their study showed no 28-day mortality difference, but they did not give the actual mortality rates (34). However, they did observe a statistically significant decrease in both the Apache II and sepsis scores from day 0 to day 4 (34). At present, this study cannot be adequately appraised, because it has not been published in a peer reviewed journal. One concern is that the study may have been ended prematurely because only 653 patients were enrolled, despite publishing an a priori sample size calculation of 800 patients (35).

**SUMMARY**

Prospective, randomized, placebo controlled clinical trials have consistently demonstrated lower incidence rates of infection, particularly pneumonia, among critically ill postsurgical patients receiving prophylaxis with IVIG. However, these studies have not shown any significant decrease in mortality, and have had variable reductions in antimicrobial use and length of ICU stay associated with IVIG therapy. Furthermore, the risk-benefit and cost effectiveness of IVIG in this setting have not been determined, and because the studies have tended to include highly selected patients, generalization beyond the study populations is difficult. Despite good evidence that IVIG reduces the incidence of infection in highly selected, critically ill postsurgical patients, its use cannot be recommended at the current time, because it has not been shown to be superior to the standard practice of antibiotic treatment of infection once it occurs. However, if the rate of infection with multidrug-resistant organisms among critically ill patients continues to rise, IVIG prophylaxis may play an increasingly important role in the management of these patients.

IVIG is a promising adjunctive treatment modality for sepsis and septic shock, but randomized trial evidence is conflicting. Small, randomized trials have shown improvement in mortality with IVIG use, and an overall significant reduction has been identified using meta-analysis. However, a generally well-designed, large, randomized trial has apparently shown no mortality benefit. It is therefore unclear whether there is a significant mortality benefit associated with the use of IVIG in sepsis and septic shock. The small trials and meta-analysis may potentially be overestimating the usefulness of IVIG, in part because of biased study designs or conduct and possibly from publication bias of trials that demonstrate significantly different results. However, the magnitude of the risk reduction observed in the best of the small studies and the meta-analysis does raise the possibility of a significant effect with IVIG treatment. On the other hand, a large, well-designed trial has shown no mortality difference, but it remains unpublished and may have been ended prematurely. At present, clinical equipoise exists for the use of IVIG in sepsis and septic shock, and further evaluation is warranted. Because of its high cost, IVIG is not currently recommended as a routine adjunctive therapy in the treatment of sepsis and septic shock.

There are a number of issues that need to be addressed in any future clinical trials of IVIG in the treatment of sepsis and septic shock. The available evidence suggests that patients with intermediate degrees of severity of septic shock appear to have the greatest benefit, and these patients should comprise the study populations. The timing and dose of IVIG also need to be optimized. Early treatment is expected to have the greatest impact by minimizing the cascade of proinflammatory cytokines. Furthermore, clinical trials, so far, have used a maximum dose of 1 g/kg IVIG that is typically given in divided doses over several days, and this may be inadequate. Based on experience with illnesses in which IVIG therapy has been proven effective, such as Kawasaki disease, a single 2 g/kg dose may be more appropriate (37). Another important consideration is that mortality may not be the best outcome measure. Organ failure and quality of life have been suggested as being more valid (38). Finally, IVIG trials should not continue to be restricted to Gram-negative infections, because this treatment particularly may be of benefit in toxigenic, Gram-positive infections. Kaul et al (39) performed a case-comparative study with group A streptococcal toxic shock syndrome patients and found a significant mortality difference of 34% of 21 cases treated with IVIG compared with 67% of 32 historical controls (P=0.02). However, these results must be interpreted with caution, because the control patients were historical, and IVIG patients were

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more likely to be treated with surgery and receive clindamycin, which may bias toward better outcomes.

CONCLUSIONS

IVIG use does prevent infection in highly selected surgical patients, but its routine use is not recommended based on a lack of documented cost effectiveness and risk-benefit. There is conflicting evidence for a mortality benefit for IVIG as a treatment adjunct for sepsis and septic shock, and further study is needed. Because of its high cost, it is not currently recommended as routine therapy. Despite a lack of good clinical trial evidence, IVIG has become a standard of care and is recommended for the treatment of streptococcal toxic shock syndrome.

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