Prehospital resuscitation
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SUMMARY
Traumatic injury is the leading cause of death in young people in the USA. Our knowledge of prehospital resuscitation is constantly evolving and is often informed by research based on military experience. A move toward balanced blood product resuscitation and away from excessive crystalloid use has led to improvements in outcomes for trauma patients. This has been facilitated by new technologies allowing more frontline use of blood products as well as use of tranexamic acid in the prehospital setting. In this article, we review current practices in prehospital resuscitation and the studies that have informed these practices.

INTRODUCTION
Traumatic injury is the leading cause of death in the USA for patients up to age 44 years.1 From the time of injury to definitive hemorrhage control, there are many opportunities to improve the ultimate outcome by minimizing physiological derangements with resuscitation as early in the course as possible. Acute traumatic coagulopathy (ATC) is the process by which the effects of severe injury combined with tissue hypoperfusion leads to a cycle of acidosis, hyperthermia, and hemodilution from crystalloid resuscitation, which has low pH, high chloride, no oxygen carrying capacity and no coagulation capacity correction. This leads to inflammatory changes, hyperfibrinolysis, endothelial dysfunction, dysfibrinogenemia, and platelet dysfunction.2 These changes exacerbate uncontrolled hemorrhage and organ dysfunction, feeding back into a self-perpetuating process which if not interrupted leads to a fatal outcome. Intervention on this lethal cycle by appropriate resuscitation should start as early as possible, ideally at the time of first contact with healthcare providers.

MILITARY EXPERIENCE WITH PREHOSPITAL RESUSCITATION
Much of our current knowledge of resuscitation in major trauma is based on military experience. The Committee on Tactical Combat Casualty Care (CoTCCC) is part of the Defense Health Agency Joint Trauma System that develops guidelines for management of injured warfighters in the field.3 Hemorrhagic shock is recognized by the presence of altered mental status in the absence of brain injury and/or weak or absent radial pulses.4 Fluid resuscitation is reserved for patients in shock, and the order of fluid priority is (1) liquid cold stored low-titer O whole blood; (2) prescreened low-titer O fresh whole blood (from a ‘walking blood bank’); (3) plasma, red blood cells (RBCs) and platelets in a 1:1:1 ratio; (4) plasma and RBCs in a 1:1 ratio; and (5) plasma or RBCs alone. Crystalloids are not listed in this protocol.

TIMING OF CRYSTALLOID RESUSCITATION
Unlike the well-established process within the military to make blood products available out of hospital, few prehospital emergency medical service (EMS) agencies have access to RBCs and plasma where the majority only have crystalloid fluid available for resuscitation. As crystalloid is not ideal for replacing blood loss, a question arises of whether crystalloid is helpful or harmful in the early stages of resuscitation. A study by Bickell et al in 1994 examined the effects of delaying fluid resuscitation until definitive hemorrhage control was obtained in hypotensive patients with penetrating torso injuries. This was a prospective study of patients with prehospital systolic blood pressure (SBP) less than 90 mm Hg in which patients were randomized to immediate standard fluid resuscitation versus no fluid. The intervention started in the field and continued until hemorrhage was controlled. Labs on arrival included hemoglobin, platelets, prothrombin time (PT) and partial thromboplastin time (PTT) were consistent with a greater degree of anemia and coagulopathy in the immediate resuscitation group. In the delayed resuscitation group, 70% survived versus 62% in the standard group (p=0.04). Whereas the intraoperative blood transfusion volume was similar in the two groups, the rate of intraoperative fluid administration was greater in the immediate group, and the total fluids and blood products administered was significantly higher in the immediate group. The delayed resuscitation group also showed a trend toward fewer complications and had a significant reduction in length of hospital stay.

VOLUME OF CRYSTALLOID RESUSCITATION
The Resuscitation Outcomes Consortium consisted of multiple trauma systems in North America (figure 1). This group performed a prospective randomized trial starting in the field where hypotensive patients were assigned to either standard resuscitation where patients received 2 L of normal saline immediately and were transitioned to TKVO (the lowest volume possible “to keep [the] vein open”) if the SBP was ≤110 mm Hg or ongoing boluses if the SBP was ≤110 mm Hg, compared with a controlled resuscitation protocol of 250 mL boluses for a SBP <70 or absence of a radial pulse. The protocol was continued until hemorrhage control was achieved or for up to 2 hours in hospital. Nineteen EMS units and 10 hospitals participated in this study in six regions of North America. Whereas no significant difference was seen in mortality in patients with penetrating trauma, patients with blunt injuries...
had significantly lower 24-hour mortality if randomized to the controlled resuscitation group.

**PREHOSPITAL BLOOD PRODUCT RESUSCITATION**

The Prehospital Air Medical Plasma (PAMPPer) trial was a multicenter cluster randomized trial comparing prehospital administration of thawed plasma compared with standard crystalloid resuscitation during air transport to trauma centers. Mortality at 30 days was significantly lower in the plasma group (almost 10% improvement). International normalized ratio (INR) was significantly lower in the plasma group and there were no differences in multiple organ failure, acute lung injury/acute respiratory distress syndrome, nosocomial infections, or allergic/transfusion reactions. This study also showed that the volume of prehospital crystalloid and RBC administration was significantly lower in the plasma group. The Control of Major Bleeding After Trauma Trial (COMBAT) trial done the same year was a single center randomized trial comparing up to two units of FFP to standard normal saline resuscitation initiated in the field in patients with SBP of <70 mm Hg or 71 mm Hg to 90 mm Hg plus HR of >108. Plasma units contained in modified bags with a large surface area were maintained frozen on ground ambulances and thawed within 3 minutes to 4 minutes using a thermal device (figure 2). This study showed no significant difference in baseline characteristics, 28-day mortality, safety outcomes or adverse events. However, the average transport time was less than 20 minutes in the COMBAT trial. In the PAMPPer trial, subgroup analysis showed no mortality benefit of plasma in patients with short transport times, whereas those with ‘prolonged’ prehospital transport times did have increased survival with plasma. However, the definition of prolonged versus short transport time was not explicitly stated in this article.

A recent review of prehospital blood transfusion found that when plasma and packed red blood cells (pRBCs) were transfused concurrently a significant improvement of long-term mortality was seen, although this was not the case with 24-hour mortality, and pRBCs transfused alone did not show any mortality benefit. Within the military population, prehospital blood product transfusion in Afghanistan was associated with greater 24-hour and 30-day survival compared with delayed or no transfusion. The majority of these patients received only pRBCs, and as such, there may be room for improvement with more balanced blood product transfusion. One way some systems have moved toward this is with increased availability of whole blood. Low-titer type O whole blood (LIOTOB) was to lead to less out of emergency department transfusion and when corrected for severity of injury and other confounders was associated with a twofold likelihood of survival. Whole blood is now the preferred resuscitative fluid of the CoTCC and despite the challenges associated with temperature requirements, cold stored LIOTOB has been effectively transported to and used in austere military environments with favorable results shown in a case series. Civilian experience with prehospital whole blood transfusion is still limited. One regional program has been established in Texas and although mortality data has not been published the cost–benefit balance seems to be favorable.
Prehospital tranexamic acid (TXA)

The CRASH-2 trial was a randomized controlled trial comparing tranexamic acid (TXA) (1 g bolus started by EMS followed by a 1 g in hospital infusion) to placebo for patients with significant hemorrhage or risk for significant hemorrhage. A small mortality benefit was seen in patients who received treatment within 3 hours of injury. The benefit was greatest the earlier TXA was given and there was a signal of benefit in TBI. CRASH 3 was a follow-up study which focused on outcomes in TBI where TXA or placebo was administered within 3 hours of injury. There was a small decrease in head injury-related death in mild to moderate head injury but not severe head injury, with all other outcomes examined being similar between groups. A recent randomized multicenter study by Rowell et al examined outcomes in patients with moderate or severe traumatic brain injury who received either 2 g of TXA in the out of hospital setting versus 1 g out of hospital plus a 1 g in hospital infusion versus placebo. The primary analysis assessing all enrolled patients did not show a difference in survival or 6 months’ neurological outcome. A planned secondary analysis on patients who had intracranial bleeding on admission head CT revealed a significant survival advantage which primarily occurred during the first 10 hours after injury. In this secondary analysis, patients who received 2 g of TXA also had improved 6 month neurological outcome. In the California Prehospital Antifibrinolytic Therapy (Cal-PAT) study, another randomized multicenter study, mortality in the group that received TXA (1 g bolus and 1 g in hospital infusion) was significantly reduced at 28 days, although there was no significant difference at 24 or 48 hours (Neeki). The mortality difference was greatest in more severely injured patients. All patients required less blood transfusion after TXA administration, and no difference in adverse events was seen between groups. Based on these data, the CoTCCC has changed their guidelines to include 2 g of TXA for patients in hemorrhagic shock or those with TBI in the field.

CONCLUSION

Prehospital resuscitation is continuously evolving. Blood product resuscitation is particularly beneficial with long transport times to prevent the initiation and progression of ATC. Crystalloid is not a preferred resuscitation for hemorrhagic shock unless blood products are not available and the patient is in shock. In this situation, only enough crystalloid to maintain organ perfusion should be given. TXA should be given as a resuscitation adjunct in the field to patients with hemorrhagic shock or TBI. Today’s special forces medics carry LTOWB, cold stored platelets, lypophilized plasma and TXA to optimize prehospital resuscitation, a combination of resources that is more than some level 1 trauma centers. More research into novel blood product resuscitation in the out of hospital setting is needed.

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