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**Massive transfusion: An update for the anesthesiologist**

Fredericks C *et al.* Massive transfusion

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**Abstract**

Exsanguination from trauma, gastrointestinal bleeding, and obstetric hemorrhage remains a major source of mortality across the planet. Continued research into resuscitation strategies and evolving technology and blood product storage has allowed for patient to undergo very large volume transfusions, even to the point of replacing a patient’s blood volume several times over. As massive transfusions have become more common, more studies have been performed delineating the exact patient population that would benefit, start- and stop-points of transfusions, complications and avoidance of the same. We discuss these points and provide information and strategies for massive transfusion.

**Key words:** Massive transfusion protocol; Exsanguinating blood loss; Blood transfusion; Trauma surgery

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**Core tip:** Recognizing the patient who requires massive transfusion early is key to the most optimal outcome. Once recognized, massive transfusion protocols (MTP) should be initiated and continued until normal physiologic parameters are reached and definitive control of bleeding is achieved. Hospitals should develop their own MTP, guided by the literature, and according to their specific needs and patient populations.

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**INTRODUCTION**

Trauma continues to be a major source of mortality, and much of this early mortality comes from exsanguination leading to death. To combat this, strategies to rapidly and effectively administer blood products and associated adjuncts to control blood loss as quickly as possible. Most deaths due to blood loss happen in the first 6 h[1]. While definitive control occurs in the operating room or interventional radiology suite, continued resuscitation is necessary to keep the patient alive during the critical first hour of the bleeding patient’s arrival to the hospital. The goal of massive transfusion is not just the replacement of intravascular volume, but the correction of trauma induced coagulopathy, in an attempt to curb further blood loss. Massive transfusion, defined as > 10 units of blood in the first 24 h can be a life-saving maneuver for a bleeding patient, but is not without complications[2]. Massive transfusion remains an area of great study throughout the critical care and trauma literature, and many hospitals now have developed their own massive transfusion protocols (MTP). This review will discuss the recent advances in massive transfusion, initiation of MTP, special populations, and complications of MTP.

**WHEN TO INITIATE MTP**

Initiation of MTP was formerly firmly under clinician gestalt. However, this intuition appears to have only a 50% predictive value in identification of patients who will need MTP[3]. Recently, clinicians developed massive resuscitation scoring systems to accurately identify patients who will ultimately require large volume blood product resuscitation upon arrival to the emergency department.

Early identification of this cohort is important before the tipping point of hemorrhage spirals into the lethal triad of coagulopathy. Cotton *et al*[4] found that both short-term and long-term survival is increased when the initiation of MTP occurs immediately in the emergency room rather than later in the operating room. Implementation of MTP guidelines both decrease mortality and the overall amount of blood use in 24 h, due to better proximal resuscitation[5].

Therefore, scoring systems for initiation of MTP must have appropriate sensitivity and also specificity to safely rule out those who will not require large volumes of blood products thus limiting unnecessary infectious exposure and saving valuable resources. These models vary in their variables of laboratory values, physical exam findings, and physiological triggers, but all have comparable predictive abilities. Application of these scoring systems is dependent on the ability for point of care testing in individual hospitals.

The German originated trauma associated severe hemorrhage score (TASH) was the first massive transfusion scoring system. It has 7 variables involving history (gender), physical findings (FAST, long-bone or pelvic fractures), vital signs (heart rate, blood pressure), and laboratory values (base deficit, hemoglobin)[6]. The probability for MTP = 1[1 + exp(4.9-0.3 × TASH)].

The Assessment of Blood Consumption score was designed to be applicable on immediate arrival to the emergency room without additional laboratory testing or need for additional calculations. It assigns one point to: Penetrating mechanism, heart rate > 90 bpm, positive FAST, or SBP < 90 mmHg. An Assessment of Blood Consumption score of greater than 2 was determined to be the appropriate trigger for MTP, with a sensitivity and specificity of 75% and 86%, respectively[7].

Cessation of MTP may more nebulous and reliant on clinician gestalt given the potentially rapidly changing current of the patients’ clinical condition in the short hours after admission. Following surgical control of bleeding, the restoration of hemodynamics, correction of acidosis and coagulopathy, and signs of sufficient end-organ perfusion (mental status, urine output) all may be used as surrogates for adequate resuscitation. Callcut *et al*[8] identified those patients at risk of ongoing hemorrhage at 6 h after admission and those patients unlikely to need additional blood products. The authors MTP score is composed of 5 variables (INR > 1.5, hemoglobin < 11 g/dL, base deficit > 6) and vital sign (SBP < 90 mmHg, temperature < 35.5 ℃). Failure of normalization of the MTP score within 3 and 6 h associated with mortality at one day and 28 d.

Compliance to MTP is vitally important, as it has been shown to negatively affect survival. In their institutional review, Bawazeer *et al*[9] found delays in 50% of activation and a 47% incidence of non-compliance with type of product given. The authors found significant differences in mortality between groups of high and medium compliance.

Despite the aforementioned algorithmic calculations for initiation and cessation of MTP, providers will often initiate MTP on the early evidence or concern for significant hemorrhage, *i.e.,* initial thoracic or pelvic radiography showing hemothorax or open book pelvic fracture, respectively. It is our suggestion to place institutional protocols to limit the number of people with the power to activate MTP to prevent overutilization by providers inexperienced with the set activation points.

**BLOOD PRODUCT RATIOS**

Many early studies in blood product ratios of fresh frozen plasma (FFP) to blood and platelet ratios were affected by survival or selection biases and mixed populations[10-12]. Most studies have focused on the risks and benefits associated with a 1:2 or 1:1 ratios of FFP to blood, although other less common ratios exist[13]. With the findings of the Pragmatic Randomized Optimal Platelet and Plasma Ratios (PROPPR) study group published in 2015, many hospitals have decided on a 1:1:1 ratio of products[14]. The PROPPR group prospectively randomized patients at 12 centers to receive 1:1:1 or 1:2:1. While 30-d mortality was similar, there was an increase in early hemostasis and decrease in deaths due to bleeding, at the expense of increased use of blood products. No increase in transfusion-related complications was noted in the 1:1:1 group[14]. The practical matter of having to thaw plasma results in many hospitals having to “catch up”, using blood first, then using plasma once it is thawed. This is disadvantageous as maintaining clotting factors and fibrinogen is important to achieving early hemostasis. A fibrinogen level lower than < 100 mg/dL substantially increases in-hospital mortality[15]. Alternatives to “catching up” include keeping a small number of units of thawed plasma and rotating it out as necessary, or using lyophilized plasma.

**ADJUNCTS TO MASSIVE TRANSFUSION**

In addition to a balanced hemostatic resuscitation (BHR) protocol multiple other pharmacologic adjuncts have been studied. Initially utilized for procedural related hemorrhage in hemophilic patients[16], tranexamic acid (TXA) was found to have application in the management of surgical bleeding. Multiple studies demonstrated a role for reduced use of blood products in elective surgery as well as the treatment of hyperfibrinolysis associated with cardiopulmonary bypass[16-19]. TXA is a lysine analog which, similar to aminocaproic acid, inhibits fibrinolysis. The antifibrinolytic effect is attributed to its competitive inhibition of the lysine binding site on plasminogen, resulting in the molecules stability and prevents further degradation of the existing clot[20,21]. In 2010 the Clinical Randomization of an Antifibrinolytic in Significant Hemorrhage (CRASH)-2 study reported the results of a 20000 patient RTC in which the effects of TXA on mortality and transfusion requirements in adult patients who were traumatically injured and in hemorrhagic shock[22]. The CRASH-2 group was able to demonstrate an improvement in all-cause mortality and mortality attributed to hemorrhage with the use of TXA within 3 h from time of injury. Subsequent studies, such as the MATTERs (Military Application of Tranexamic Acid in Trauma Emergency Resuscitation) and PED-TRAX (Pediatric Trauma and Tranexamic Acid), demonstrated a survival advantage when TXA was administered[23,24]. In the MATTERs study there was an observed lower unadjusted mortality rate, increased odds of survival and lower rate of coagulopathy[23]. With PED-TRAX, the first study to look at children independently, there was an observed decreased mortality among all patients (OR = 0.3; *P* = 0.03) who received TXA[24]. Interestingly both populations of the MATTERs and PED-TRAX studies who received TXA were more severely injured than those who did not yet there remained an observed survival benefit.

Regarding thromboembolic events, there was an observed increase in pulmonary embolus and deep venous thrombosis in the MATTERs study but these results were not replicated in the PED-TRAX study[23,24]. While there appears a clear benefit to use of TXA in the bleeding trauma population, several critiques have arisen specifically regarding the CRASH-2 study. These include the applicability and predictability, lack of measurement of severity of injury and the application of TXA use in traumatic brain injury[25]. Several upcoming studies like CRASH-3 and the Prehospital Antifibrinolytics for Traumatic Coagulopathy and Hemorrhage Trial will hopefully elucidate a more specified role for TXA, but what we do know is that TXA provides some benefit to the bleeding trauma patient, and is included in many transfusion protocols[26,27].

The use of supplemental fibrinogen, Recombinant Factor VII (rFVIIa), and prothrombin complex (PCC) as adjuncts to BHR have also been explored. Several nonrandomized trials have evaluated the use fibrinogen in the trauma patient[28-30]. In each of these studies there was a reduced requirement for blood product transfusion and associated reduced potential for multi-organ failure. While there has been shown a benefit, no prospect randomized trials have been reported to date. rFVIIa has also been examined for use in massive transfusion. Multiple retrospective studies have elevated the added effect of rFVIIa on volume of transfusion, mortality and organ failure with mixed results[31-34]. To date, only two randomized controlled trials have been completed[35,36]. Unfortunately, neither study revealed any mortality benefit with the addition of Factor VII though there was observed reduction in amount of blood products transfused, furthermore a Cochrane review was unable to support the use of Factor VII for use in traumatic hemorrhage[37]. PCC has also been proposed for use in BHR. Initially utilized in the treatment of hemophilia, it is the standard reversal agent for vitamin K antagonists and has been utilized in BHR[38]. Several well-constructed reviews and retrospective studies have shown efficaciousness of PCC in addressing vitamin K antagonistic bleeding in trauma, but no clear role in it use in BHR has been defined[39-41]. Hannon *et al*[42] in an animal based model of traumatic coagulopathy failed to show any benefit on blood volume lost when PCC was utilized and there is currently a lack of prospective randomized controlled trials to guide evidence supported use. Although promising, additional prospective study is needed of concentrated fibrinogen or prothrombin complex concentrate before they can be recommended[43].

While a balanced hemostatic resuscitation is the cornerstone of addressing the bleeding trauma patient, there remains much work to be done in the investigation of the multiple potential and available adjuncts before the optimal strategy is determined.

**BLOOD ADMINISTRATION METHODS**

Speed of transfusion is important during MTP especially when exsanguination occurs at a rate greater than transfusion. Speed is augmented by optimal vascular access and utilizing Pouiselle’s law, which states flow is directly related to width and indirectly related to length of the catheter. Pressurized tubing or specialized pressurized rapid transfusers can augment flow. Rapid transfuser units can often also warm blood at the same time, providing an additional benefit against hypothermia.

Blood and fluid warming is important as to not to exacerbate hypothermia. Barthel *et al*[44] suggest warmed fluids cannot induce hyperthermia, but can limit additional heat loss. However, adjunctive techniques of warming including forced air devices, blankets, and high operating room temperatures are important. Hypothermia is exceedingly dangerous; in patients undergoing MTP, a cutoff of less than 35 ℃ as the lowest recorded temperature in the first 24 h portended increases in mortality[45].

**SPECIAL POPULATIONS**

***Elderly patients***

The elderly patient represents a unique challenge amongst the trauma population at base line. Underlying cardiovascular disease and decreased functional status can complicate resuscitation. Data from the PROMMTT and PROPPR had median ages for 37 and 34 years respectively, which questions the applicability of such practices to an elderly population[14,46]. The Trauma Outcomes Group demonstrated that age is an independent predictor of mortality in the massively transfused patient[2]. In theory the decreased need for crystalloid volume would lead to a decreased incidence of circulatory over load in the elderly population. Although no subgroup analysis of the PROMMTT or PROPPER studies exists and no prospective institutional studies exist to examine age in a massive transfusion protocol. In a retrospective analysis of 14 elderly (> 60 years) compared to 52 non-elderly patients, Murry *et al*[47] demonstrated similar, mortality rates (50% *vs* 53%). This is a limited study with high mortality in both arms, but otherwise similar patient characteristics on arrival including GCS, ISS and starting hematocrits. Mitra *et al*[48] retrospectively compared patients > 65 years *vs* a younger cohort and demonstrated that while mortality was higher (39% elderly *vs* 21% young) a significant number of elderly patients survived to discharge. Patient characteristics that were associated with mortality among the elderly patients included an increased systolic blood pressure (OR = 1.02), a pre hospital GCS < 8 (0.73), and acute traumatic coagulopathy (11.75). Limited data exists for the use of massive transfusions in the elderly trauma patient. Small series would suggest similar mortality to cohorts from the same institutions, further work to explore coagulopathic complications is needed. Additionally, considerations for age, frailty and cardiovascular function should be investigate and possibly included in future targeted massive transfusion protocols.

***Pediatric patient***

The pediatric patient population also represents unique challenges to hemorrhagic shock and acute traumatic coagulopathy. The physiologic reserve in children is robust and includes the ability to maintain normal blood pressures until 20% of blood volume loss[49]. Which make application standard triggers used in adults difficult to apply. Resuscitation in the pediatric population is defined on volume (mL) per body weight (kg) and is no different in acute hemorrhage. Children younger than 3 mo are estimated to 90 mL/kg of blood volume and those older than 3 mo are estimated at 70 mL/kg[50]. Some measures do apply: High ISS scores, shock, high base deficit and increased INR > 1.5 are associated with increased mortality in the pediatric population[51]. In an effort to better define the volume of blood loss needed to qualify for a pediatric MTP, Neff *et al*[52] used the DOD trauma registry and identified all pediatric patients, those to greater than 40 mL/kg of blood loss in the initial 24 h were more often in shock, hypothermic, coagulopathic and thrombocytopenic at time of presentation[52,53]. This gives a reasonable initial starting point for a volume trigger for pediatric MTP. The same principles apply including balanced transfusion strategies and a restrictive use of crystalloids. That stated there are only two small prospective trials on the use of MTP in pediatrics. Hendrickson *et al*[54] described the initiation of protocol involving a fixed ratio of products based on body weight. They included 102 patients, and succeeded in transfusing at nearly a 1:1 FFP to RBC ratio, although no statistic improvement was seen in mortality (38% pre *vs* 23% post *P* = 0.035) after taking into account severity of injury. In this study only 50% of patients required a massive volume of blood (> 70 mL/kg) to be transfused this may reflect our inability to identify pediatric patient in need and not the benefit of MTP. Chidester *et al*[55] also applied the MTP principles to the pediatric population; they included 55 patients with transfusion ratios at 1:3 and similarly didn’t see an improvement in mortality. They did note fewer thromboembolic complications with the MTP group (4 events *vs* 0 events).

***Obstetric patients***

Post-partum hemorrhage (PPH) is a major cause of up to 25% of pregnancy related deaths[56]. Physiologic changes in pregnancy including an increase in red blood cell mass (25%) and a greater increase in plasma volume (50%) allow of hemodynamic stability during the birthing process. Although changes in coagulation factors are not balanced and the relative increase in fibrinogen and factors VII, VIII and IX lead to a relative hypercoagulable state[57]. These physiologic changes make the resuscitation of such patients uniquely challenging. Primary treatment of postpartum hemorrhage includes surgery and uterotonic agents. Some investigators are starting to apply balanced and goal directed resuscitation to PPH[58]. Adjuncts such as thromboelastography allow for a real-time assessment of coagulation and fibrinolysis and have allowed for targeted treatment[59,60]. Although familiarity with the normal baseline changes in the tests are key for interpretation[59]. Despite these initial advancements in PPH care, the lack of prospective data has led to limited improvement in the national management guidelines. Dahlke *et al*[61] compared management guidelines from the 4 large nations obstetrics committees and only one guideline includes mention of blood bank notification and none describe a balanced or targeted transfusion practice. Further prospective analysis is required for the use of massive transfusion in the obstetric patient.

**COMPLICATIONS OF MASSIVE TRANSFUSION**

Administration of liters of inflammatory, immunomodulatory, and potentially infectious fluids into a patient already in hemorrhagic shock can assist in explanation of the common morbidities of MTP. They include: acute respiratory distress syndrome (ARDS), transmission of viral and bacterial infection, abdominal compartment syndrome (ACS) and electrolyte abnormalities.

***Lung injury***

Massive resuscitation makes the lungs susceptible to the spectrum of lung injury *via* volume or immune-mediated mechanisms. For those undergoing MTP, Moss *et al*[62] found that 21% of patients will develop ARDS. The risks appear to increase with the amount of blood transfused and the mechanism of injury. Silverboard *et al*[63] demonstrated in a prospective cohort of 102 patients, development of ARDS was found at 5 or 10 units of PRBCs transfused for blunt or penetrating trauma respectively.

***Infectious risk***

MTP places patients at risk for both viral and bacterial infections. In the United States, the estimated risk for HIV is 1 in 2135000. The greatest risk is for Hepatitis B at 1 in 277000[64]. Patients are more susceptible to bacterial infections from platelets because of their relatively warm storage requirements (20 ℃), with an estimated risk of 1 in 5000 compared to 1 in 38500 in RBC[64]. The most common bacteria transmitted are gram-positive aerobic organisms[65].

***Compartment syndrome***

The incidence of intraabdominal hypertension or abdominal compartment syndrome appears to be declining with limitation and replacement of crystalloid with balanced blood products strategies. Joseph *et al*[57] showed a decrease in ACS from 7.4% to 0% with corresponding reduction in crystalloid from 12.8 to 6.6 L.

***Electrolytes***

Two common electrolyte abnormalities that occur in MTP are hypocalcemia, caused by the preservative citrate and hyperkalemia. Aboudara *et al*[66] show that after transfusion of 7 units of PRBCs, the patient is at risk for hyperkalemia. Furthermore, in the pediatric population, there is an association between hyperkalemia during rapid blood transfusion and cardiac arrest.

**CONCLUSION**

The finer points of massive transfusion, such as ratios of blood products, adjuncts, and transfusion triggers and endpoints will continue to be studied and debated in the literature for years to come. However, it cannot be argued that a massive transfusion protocol saves lives. A defined, hospital-specific MTP allows trained providers to recognize patients at risk of high-volume blood loss early, initiation of massive transfusion quickly, and has specific stop points to limit over transfusion[67]. We recommend that all centers that take care of critically ill patients of all varieties should evaluate the literature and develop their own protocol. We have included our protocol for guidance (Figure 1).

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**Figure 1 Our hospital-specific protocol for massive transfusion.**