Prehospital Blood Transfusion (CPG ID:82)
The CPG provides a brief summary of the scientific literature for prehospital blood use and essential instructions on resuscitation procedures using blood products.

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Publication Date: 30 Oct 2020

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PURPOSE

This Clinical Practice Guideline (CPG) provides a brief summary of the scientific literature for prehospital blood use, with an emphasis on the en route care environment. Updates include the importance of calcium administration to counteract the deleterious effects of hypocalcemia, minimal to no use of crystalloid, and stresses the importance of involved and educated en route care medical directors alongside at a competent prehospital and en route care providers (See Table 1.) With the paradigm shift to use FDA-approved cold stored low titer group O whole blood (CS-LTOWB) along with the operational need for continued use of walking blood banks (WBB) and point of injury (POI) transfusion, there must be focused, deliberate training incorporating the different whole blood options. Appropriate supervision of autologous blood transfusion training is important for execution of this task in support of deployed combat operations as well as other operations in which traumatic injuries will occur. Command emphasis on the importance of this effort as well as appropriate logistical support are essential elements of a prehospital blood program as part of a prehospital/en route combat casualty care system.

Table 1. Five Levels of Clinical Competence

<table>
<thead>
<tr>
<th>Skill Level</th>
<th>Description</th>
<th>Trauma-related Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Novice</td>
<td>The novice has no experience in the environment in which they are expected to perform.</td>
<td>Administrator or technician who has never worked in a trauma center.</td>
</tr>
<tr>
<td>Advanced Beginner</td>
<td>The advanced beginner demonstrates marginally acceptable performance and has enough experience to note recurrent meaningful situational components.</td>
<td>Medic that has had didactic trauma training but no clinical trauma experience.</td>
</tr>
<tr>
<td>Competent</td>
<td>Competence is achieved when one begins to see one's actions in terms of long-range goals or plans. This individual demonstrates efficiency, coordination, and confidence in his/her actions.</td>
<td>Board eligible/certified physician, newly trained nurse, but has only rotated as a learner at a trauma center.</td>
</tr>
<tr>
<td>Proficient</td>
<td>The proficient individual perceives situations holistically and possesses the experience to understand what to expect in a given situation.</td>
<td>Board eligible/certified physician or newly assigned nurse with limited but recent prior exposure to trauma, starting their career at a high volume and best quality Level I trauma center.</td>
</tr>
<tr>
<td>Expert</td>
<td>The expert has an intuitive and deep understanding of the total situation and is able to deliver complex medical care under highly stressful circumstances.</td>
<td>Trauma nurse coordinator, Board certified Emergency Physician or fellowship trained trauma surgeon/intensivist with years of experience at a high volume and best quality Level I trauma center.</td>
</tr>
</tbody>
</table>

Sources: Adapted from Benner, 1982; Dreyfus, 1981; Dreyfus and Dreyfus, 1980.
INTRODUCTION

Early administration of blood products to the trauma patient in extremis is the standard in combat casualty care and becoming more common as part of a strategy of civilian prehospital critical care.\(^1\)\(^-\)\(^7\) Although there is some dissent regarding the level of evidence for and benefits of prehospital blood transfusion in the civilian literature, much of the data informing that discussion precedes the use of whole blood and involves a patient population dissimilar from the military.\(^8\)\(^,\)\(^9\) Civilian Emergency Medical Services systems internationally\(^10\),\(^11\) and in Texas, North Carolina and several other states have initiated whole blood transfusions in the prehospital environment. At least one civilian tactical law enforcement team has initiated a Low-Titer O Fresh Whole Blood program.\(^12\)

The main military literature covering this topic is summarized in Table 2 below.

### Table 2. Military literature on prehospital transfusion

<table>
<thead>
<tr>
<th>Article</th>
<th>Setting</th>
<th>Inclusion</th>
<th>Findings</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shackelford SA, Del Junco DJ, Powell-Dunford N, et al. (\text{Association of Prehospital Blood Product Transfusion During Medical Evacuation of Combat Casualties in Afghanistan With Acute and 30-Day Survival.}) JAMA. 2017;318(16):1581-91.</td>
<td>Mostly Military air ambulance evacuations</td>
<td>1) Traumatic limb amputation at or above the knee or elbow or 2) Shock defined as a systolic blood pressure of less than 90 mm Hg or a heart rate greater than 120 beats per minute.</td>
<td>Adjusted hazard ratio for mortality associated with prehospital transfusion was 0.26(95%CI, 0.08 to 0.84,(P=0.02)) over 24 hours (3 deaths among 54 recipients vs 67 deaths among 332 matched nonrecipients) and 0.39(95%CI,0.16 to 0.92,(P=0.03)) over 30 days(6 vs 76 deaths, respectively)</td>
<td></td>
</tr>
<tr>
<td>Malsby RF, 3rd, Quesada J, Powell-Dunford N, et al. (\text{Prehospital blood product transfusion by U.S. army MEDEVAC during combat operations in Afghanistan: a process improvement initiative.}) Mil Med. 2013;178(7):785-91.</td>
<td>Military air ambulance in Afghanistan</td>
<td>Traumatic injury (chest/abd and single or multiple amputations) with 1) systolic blood pressure (SBP) &lt;90 mmHg 2) heart rate (HR) &gt;120 bpm 3) Oxygen Saturation ((SaO_2)) &lt;90%.</td>
<td>Feasibility study showed: 15 patients given 19 units no adverse reactions and no instances of blood product temperature outside of the accepted range</td>
<td></td>
</tr>
<tr>
<td>O’Reilly DJ, Morrison JJ, Jansen JO, et al. (\text{Prehospital blood transfusion in the en route management of severe combat trauma: a matched cohort study.}) J Trauma Acute Care Surg. 2014;77 (3 Suppl 2):S114-20.</td>
<td>Regional Command South in AFG by MERT</td>
<td>MERT Physician order based upon injury severity and blood product availability</td>
<td>Matched cohort study of 1,592 patients in Afghanistan transported by a prehospital intensivist team found mortality significantly lower in those patients who received prehospital blood transfusions than those who did not (8.2% vs 19.6%, (p&lt;0.001))</td>
<td>Confounded by different rates of invasive procedures</td>
</tr>
</tbody>
</table>
(HR) >120 bpm or Oxygen Saturation (SaO2) <90%. Following the fifth transfused unit, oxygen saturation was excluded as a transfusion trigger and multiple amputations (with at least one proximal amputation) were included as transfusion triggers. There were no adverse reactions and no instances of blood product temperature outside of the accepted range.

Advanced clinical care during patient transport has demonstrated improved outcomes. A growing body of evidence suggests that early blood transfusion, both at point of injury and en route to a surgical capability improves survival in severely injured patients with acute blood loss. Rapid initiation of blood transfusion among wounded personnel in Afghanistan was evaluated among 502 patients. Patients who had been transported via MEDEVAC; who had suffered a traumatic amputation at/above the knee/elbow; had SBP <90 mmHg or HR >120 bpm; and who had received blood products during transportation, or, shortly after arrival in the hospital, were compared to those who did not receive blood. Those who received blood were significantly less likely to die at 24 hours (adjusted hazard ratio for mortality 0.26 [95% CI 0.08-0.84, p=0.02]) as well as at 30 days (adjusted hazard ratio for mortality 0.39 [95% CI 0.16-0.92, p=0.03]). This study underscores the importance of early transfusion. The en route care providers must continue (or in some cases start) transfusion as soon as the casualty is in the aircraft. Blood products that have started transfusion at POI can be continued during MEDEVAC.

A matched cohort study of 1,592 patients in Afghanistan transported by a en route care intensivist team found mortality to be significantly lower in those patients who received prehospital blood transfusions than those who did not (8.2% vs 19.6%, p<0.001). Several confounding variables make these data difficult to interpret (e.g., differing rates of prehospital rapid sequence induction, administration of pharmacologic adjuncts and varied transportation times). Nevertheless, it is likely that an aggressive approach to prehospital critical care including blood transfusion was part of an overall strategy of damage control resuscitation that improved survival in severely injured patients.

**PREHOSPITAL PRINCIPLES OF RESUSCITATION AND TRANSFUSION**

- Rapid recognition of life-threatening hemorrhagic shock
  - Point of care devices if available: international normalized ratio; lactate level may be of value.
- Prevent hypothermia
- Hemorrhage control with mechanical hemostatic adjuncts:
  - Tourniquet/junctional tourniquet
  - Pressure dressings/thrombin and fibrin impregnated gauze
- Hemostatic Resuscitation
  - Whole blood (WB) is optimal
    1. FDA approved CS-LTOWB
    2. Low titer group O whole blood (LTOWB)
    3. Type O-WB (non-tittered)
    4. Component therapy with plasma (dried, liquid, or, thawed), red blood cells (RBCs), and platelets in 1:1:1 ratio
- Avoid hypocalcemia/Calcium replacement
In prolonged evacuations, empiric calcium administration for every 4-6 units of RBCs or WB
- Avoid crystalloid resuscitation
- Tranexamic Acid administration if less than 3 hours from time of injury
- Consider source of Freeze Dried Plasma if available

**NOTE:** Conventional goal is systolic blood pressure >90mmHg. Recent concept indicates a higher goal of 90-110mmHg due to shift towards blood-based resuscitation and concern for prolonged hypoperfusion especially for patients with long transport times.

**CRITERIA FOR TRANSFUSION**

Prehospital blood transfusion is predicated upon clearly defined criteria for the use of this valuable resource. Several high performing military and civilian prehospital medical teams use prehospital blood transfusion according to protocolized guidelines. The U.S. Army Ranger Regiment was an early adopter of prehospital blood transfusion and uses the following criteria for transfusion: signs and symptoms of hemorrhagic shock; OR 1+ amputation; blunt/penetrating trauma (junctional/abdominal/thoracic); OR pelvic fracture; OR SBP <100; OR lactate >5; pulse >100. In Australia, the Greater Sydney Area Helicopter Emergency Medical Services (GSA-HEMS) is an intensivist-based prehospital critical care service that has extensive experience in prehospital blood transfusion. In the prehospital environment, they transfuse blood if there is “persistent haemorrhagic shock despite hemorrhage control measures after crystalloid infusion.” During interhospital transport GSA-HEMS transfuses blood if there is “persistent hemorrhagic shock where there is limited or no access to cross-matched blood and [an] ongoing requirement for transfusion.” The Australian Queensland Ambulance Service Trauma Response Team administers blood products in the prehospital environment and a retrospective review of cases was published in 2014 demonstrating benefit in appropriate clinical situations. In 2014, proposed criteria for prehospital blood products in combat casualties were refined based on data from Afghanistan and updated in 2020 using a larger data set from the entire DoD Trauma Registry (DoDTR). A case series reviewing the benefits of early whole blood administration in combat casualties was published in 2016.

Development of transfusion criteria is an important consideration in establishment of a prehospital transfusion program. In the civilian trauma center experience, 596 trauma patients were evaluated using an assessment of blood consumption (ABC) score. Patients received one point for any of the following: penetrating mechanism; positive focused assessment sonography for trauma (FAST); systolic blood pressure less than 90 mmHg; heart rate greater than or equal to 120 beats per minute. An ABC score of two or greater predicted the need for massive blood transfusion with 75% sensitivity and 86% specificity. This investigation demonstrated the utility of a quick, non-laboratory based tool to predict the need for blood transfusion.

Careful attention to transfusion triggers can limit wastage of this important resource in a potentially resource constrained environment. Furthermore, it is important to note that blood product usage can be optimized when hemorrhage control is undertaken simultaneously. Ensure that hemorrhage control efforts are properly employed in accordance with Tactical Combat Casualty Care Guidelines. When administering blood, be certain to alert the receiving facility that the patient is requiring blood transfusion, noting any potential need for massive transfusion.
PROCEDURE

Certain special considerations for blood transfusion have been enacted in other Department of Defense transfusion algorithms. Rapid transfusion of blood can cause sheering of RBCs and should be avoided if possible. When possible, low titer group O whole blood (LTOWB) should be administered as the blood product of choice. LTOWB has been screened for Anti-A and Anti-B antibodies. The titer of these antibodies is low enough to represent minimal risk of clinical consequences, and may be considered a universal donor. LTOWB will be exclusively drawn from sites approved by the Armed Services Blood Program (ASBP), distributed in theater via the ASBP blood distribution system and fully tested in accordance with FDA guidelines.

Blood products should be transfused in a plasma:platelet:RBC ratio of 1:1:1. If platelets are not available, then plasma:RBC should be transfused in a 1:1 ratio. If that is not possible, then reconstituted dried plasma, liquid plasma or thawed plasma alone or RBCs alone should be transfused.

If available, type O negative RBCs should be used preferentially for females of childbearing years. If it is necessary to administer O positive blood to a woman of childbearing years with O negative blood, then consider Rhogam therapy to decrease the risk to subsequent pregnancies. Please note though that resuscitation and prevention of exsanguination takes precedence of potential future pregnancy complications. Pediatric blood transfusion should be based on clinical signs of shock rather than predefined vital signs, since vital signs can vary considerably between age groups in pediatric patients.

Blood plasma contains important clotting factors and can be transfused rapidly. Thawed plasma has a shelf life of 5 days and may not be available for prehospital missions. “Never frozen” liquid plasma has a shelf life of 26 days and may be more readily available. Check with the issuing facility for availability of thawed or liquid (i.e. never frozen) plasma. All blood products must be transfused immediately after removal from the storage container. If transfusion is delayed, then blood products that have been removed from storage and exceeded proper temperature will be returned to the issuing facility per local policy. Ensure that all blood products issues have a Safe-T-VUE (NSN 6515-08-T00-3056) attached and activated for temperature monitoring. (See Appendix E).

QUALITY CONTROL

Proper storage, transport and administration of blood and blood products is addressed the following Joint Trauma System CPGs: Damage Control Resuscitation, Frozen and Deglycerolized Red Blood Cells and Whole Blood Transfusions.

Ensure that the blood product container (NSN 6530-01-505-530I; OCP/5306; Desert) is properly charged and maintained prior to loading blood products. Ensure that all blood products have a Safe-T-VUE attached and activated for temperature monitoring. Ensure that thawed plasma is at a refrigerated temperature (1-6 degrees C) prior to placement of Safe-T-VUE. Appropriately document issuance and use of blood products per local protocol, including documentation in Theater Medical Data Store (TMDS) inventory.

Blood products carried outside of a medical treatment facility (MTF) and/or laboratory will be contained in an approved storage container for a maximum of 48 hours. Once loaded and sealed, a container will remain closed and intact until used for patient care or returned to the issuing facility. Units will inspect blood product containers and document appropriate quality metrics (e.g., intact, no leaking) every 24 hours.
TRANSFUSION REACTION

The rate of non-infectious transfusion reaction is low. In a recent evaluation of 4,857 transfusion episodes approximately 1% were associated with a serious reaction. Transfusion-associated circulatory overload was most common (~1%); while transfusion-related acute lung injury, anaphylactic reaction and hypotensive reactions all occurred in less than 0.1% of transfusions. Other investigators evaluated emergency transfusion of 5,203 units of type-O RBC to 480 trauma patients and found that no acute hemolytic transfusion reactions occurred.

In the unlikely event of a prehospital transfusion reaction, immediately stop the transfusion. In cases of anaphylaxis, administer 0.3 mL of 1:1000 epinephrine intramuscularly (IM), administer diphenhydramine 25 mg IV or IM and consider methylprednisolone (the FDA approved dose of methylprednisolone is 10 – 40 mg IV over several minutes, with subsequent doses being determined by clinical response). Maintain the patient’s airway as needed and administer IV fluids in cases of hypotension. In cases of acute hemolytic reaction, administer diphenhydramine 25 mg IV or IM, and consider osmotic diuresis with mannitol 20% 20 gm or 3% sodium chloride 250 mL.

HYPOCALCEMIA

Calcium administration should be considered in all patients undergoing en route care blood transfusion. The citrate preservative in blood products can chelate calcium and contribute to hypotension in patients who have received blood products. In 352 patients who were critically bleeding and required massive transfusion, investigators found that hypocalcemia worsened mortality. Patients generally had low ionized calcium concentration (mean = 0.88mmol/L) and had higher odds of mortality (Odds Ratio = 1.25, 95% Confidence Interval 1.04-1.52; P = 0.02). Admission hypocalcemia is further associated with mortality. Calcium concentration was measured in 591 trauma patients before administration of blood products. Those with an ionized calcium < 1 experienced higher mortality (15.5%) compared to those with ionized calcium levels >1 (mortality = 8.7%, p = 0.036).

Trauma patients are often hypocalcemic without regard to transfusion status, further suggesting that calcium administration may be beneficial. In 212 trauma patients with a mean Injury Severity Score (ISS) of 34; 64% were found to be hypocalcemic (with calcium < 1.15 mmol/L) and 10% were severely hypocalcemic (<0.9mmol/L). Hypocalcemia is common in massive transfusion and associated with increased mortality. Although the mechanism of this effect is not completely understood, it is thought that acidemia related to tissue hypoperfusion may play a role. Administration of one gram of calcium (30 ml of 10% calcium gluconate or 10 ml of 10% calcium chloride) IV/IO before, during (using a secondary access point) or immediately after the first unit of blood product. Re-dose one gram of calcium IV/IO after every 4 units of blood products.

TRAINING

Execution of prehospital and en route care blood transfusions requires sufficient training to adequately prepare. Conducting the most realistic training possible has been shown to improve in-hospital trauma team performance as well as the widely held belief of military training in general. Whole blood training should be viewed similarly. Coordinated and deliberate autologous blood labs with appropriate oversight can safely achieve the highest degree of realism with relatively low cost. Armed services blood bank policies for donor allowance in autologous blood lab participants is still under discussion.
PERFORMANCE IMPROVEMENT (PI) MONITORING

POPULATION OF INTEREST

1. All patients who receive blood product transfusion within 3 hours of injury
2. All patients who meet criteria for blood transfusion (severe traumatic injury: ISS ≥16 and ≥ 2 body regions injured with AIS severity ≥ 2 AND SBP < 100 OR HR > 100 OR hematocrit < 32% OR pH <7.25) within 3 hours of injury

INTENT (OUTCOME)

1. LTOWB is used for prehospital resuscitation of casualties with life-threatening injuries and hemodynamic instability (HR > 100 or SBP < 100).
2. For the population of interest, the first resuscitation fluid given after injury is a blood product, ideally cold-stored LTOWB.
3. Administration of whole blood and blood products to appropriate combat casualties in the prehospital and en route care environment.

PERFORMANCE/ADHERENCE METRICS

1. The number and percentage of patients in the population of interest who receive WB transfusion prior to arrival at first role of care.
2. Number and percentage of patients in population of interest who received a blood product as the first resuscitation fluid.
3. Number and percentage of patients in population of interest who received cold-stored LTOWB as the first resuscitation fluid.
4. Indication identified.
5. Pre and post vital signs measured.
6. Calcium administration documented.
7. Appropriate information collected for donor and recipient exposure monitoring.

DATA SOURCES

- Patient Record (DD1380/DA4700 OP7)
- DoDTR

SYSTEM REPORTING & FREQUENCY

The above constitutes the minimum criteria for PI monitoring of this CPG. System reporting frequency will be performed annually; additional PI monitoring and system reporting may be performed as needed.

The system review and data analysis will be performed by the JTS Chief and the JTS PI Improvement Branch.
RESPONSIBILITIES

It is the Unit Medical Director's responsibility to ensure familiarity, training, appropriate compliance and PI monitoring at the local level with this CPG. Coordination with the Regional Medical Director and Trauma Director is also essential in ensuring full spectrum compliance and PI monitoring along with clear communication with the Combatant Command Trauma System and Joint Trauma System.

REFERENCES


27. Via DK. Urgent resuscitation using blood products during tactical evacuation from point of injury. CENTCOM Clinical Operating Protocols 2016;CCOP-01.


APPENDIX A. PREHOSPITAL BLOOD TRANSFUSION PROCEDURES

BEFORE

Criteria for Blood Transfusion:

- HR >100 bpm; or
- SBP <100 mmHg or no radial pulse; or
- Altered mental status with signs/symptoms hemorrhagic shock; or
- Penetrating trauma to chest/abdomen, junctional injuries; or
- Pelvic fracture; or
- Any above knee amputation or multiple amputations (regardless of vital signs)

DURING

1. Obtain IV/IO access
   a. Verify blood product identification number
   b. Connect IV tubing with filter to blood
   c. Transfuse as rapidly as tolerated

2. Monitor for signs of transfusion reaction:
   Hypotension, flushed face, wheezing, fever, rigors, flank pain
   - Anaphylaxis = Epinephrine 0.3 mL of 1:1000 IM; Diphenhydramine 25 mg IV/IM;
     Maintain airway; Administer IV fluids prn; Consider Methylprednisolone
   - Acute hemolytic reaction = Diphenhydramine 25 mg IV/IM; Consider osmotic diuresis
     with 20 gm mannitol 20% or 250 mL 3% NaCl

AFTER

Resupply from pre-designated source: Consider 1 gm CaCl2 IV

Abbreviations: Heart rate (HR), Beats per minute (bpm), Systolic blood pressure (SBP), Intravenous (IV), Intramuscular (IM), As needed (prn), Gram (gm), Sodium chloride (NaCl), Calcium chloride (CaCl2)
APPENDIX B: RESUSCITATION USING BLOOD PRODUCTS DURING TACEVAC


PURPOSE

To provide essential instructions on urgent/life-saving resuscitation procedures using blood products during tactical evacuation (refers to both casualty evacuation and medical evacuation) from the point of injury (POI) for casualties suffering major blood loss/massive hemorrhage. Referred to as the Vampire Program. This guideline supports the Joint Trauma System (JTS) Damage Control Resuscitation Clinical Practice Guideline (CPG), Whole Blood Transfusion CPG, and the Tactical Combat Casualty Care (TCCC) Guidelines.

Guidance applies to medical and non-medical personnel (e.g., flight medic, crew chief, registered nurse, enlisted medical personnel, physician, nurse practitioner, or, physician assistant), assigned/attached or allocated to perform tactical medical response (TCCC) and evacuation (CASEVAC and MEDEVAC) duties that involve direct or indirect patient care.

FIELD INDICATIONS FOR TRANSFUSION DURING TACTICAL EVACUATION

The following are indications for transfusion in the presence of severe traumatic injury:

- Systolic BP <100 or absence of radial pulse; or
- Heart rate >100; or
- Any above knee amputation or double/triple/quadruple amputation (regardless of vital sign indication).

**WARNING:** The amputation patterns above are the only traumatic injuries that constitute a stand-alone immediate field indicator for transfusion that requires no confirmation with vital sign parameters.

**CAUTION:** Control external bleeding before or simultaneously with initiation of blood product transfusion.

Traumatic Arrest: patient with exsanguination who had signs of life when received from ground forces and has since become pulseless should receive immediate transfusion (transfusion is more important than chest compressions in cases of exsanguination and should take priority).

Traumatic injuries where early blood transfusions are most likely to be needed:

- Penetrating thoracic/abdominal/junctional (junctional includes axilla/inguinal/cervical) injury.
- Pelvic fracture.
- Multiple injuries.
- Proximal amputations (above knee or elbow). Amputation is defined as any severe trauma to a limb that involves complete or partial loss of the limb (this includes limbs that are severely mangled but not completely severed).
Initiate transfusion with 1 unit of blood product. Give additional units if clinically indicated. Avoid resuscitation with crystalloid which may increase bleeding, particularly from non-compressible torso hemorrhage.

Refer to Appendix C for list of clinical indicators for hemorrhagic shock.

**PROCEDURE**

**Blood Component Therapy Approved for Transfusion during Tactical Evacuation**

Red blood cells (RBCs) increase the recipient’s oxygen-carrying capacity by increasing the mass of circulating red cells. Plasma and platelets work together to improve blood clot formation and clot stability. On average a unit of whole blood (WB) contains a volume of 500-600 mL and a unit of RBC’s contains a volume of 300-400 mL. In an exsanguinating patient, a unit of blood can be given quickly.

Ensure good blood flow through IV or IO access before initiating transfusion. Refer to Appendix D for Transfusion Procedures and Appendix F for Pearls for Transfusion.

**CAUTION: Rapid infusion against resistance can cause mechanical shearing of RBCs and should be avoided.**

1. Blood products will be administered in the following priority depending on availability and according to TCCC Guidelines:
   a. Cold Stored Low titer Group O whole blood (LTOWB), or, if not available
   b. Fresh Low titer Group O whole blood (LTOWB), or, if not available
   c. Plasma, RBCs, and platelets in a 1:1:1 ratio, or, if not available
   d. Plasma and RBCs in 1:1 ratio, or, if not available
   e. Reconstituted dried plasma, liquid plasma, or, thawed plasma alone or RBCs alone, or, if not available
   f. Fresh group O (typed by Eldon card or equivalent) whole blood from unknown titer walking blood bank

**NOTE:** LTOWB has been screened for anti-A and anti-B antibodies; these units contained a low titer of anti-A and anti-B and are therefore considered a universal donor product that may be given to recipients of any blood type with a minimum risk for a minor ABO incompatibility (typically minor and most often subclinical clinical consequences). The whole blood supplied to MEDEVAC units will be exclusively drawn in the United States from the ASBP-approved sites and distributed in theater by the ASBP blood distribution system. The LTOWB units will be fully tested following FDA current guidelines.

2. POS (either low titer Group O WB or Type O RBCs) is the standard for transfusion during evacuation.
NOTE: Patients requiring blood can safely receive un-cross matched low titer Group O WB or Type O RBCs until type-specific products are available.

3. If available, use O NEG on females of childbearing potential age <50 years old. Inform receiving facility regarding female given O POS blood for documentation in the medical record. If a minimal amount (just a few milliliters) is given, consider Rhogam therapy. The immunologic consequences of administration of an entire unit of O POS whole blood or RBC to an O NEG female of childbearing potential cannot safely be reversed with Rhogam. Treatment of exsanguination takes precedence over potential future pregnancy outcomes.

CAUTION: WB collected in theater will NOT be supplied for use onboard MEDEVAC aircraft.

Plasma is recognized as an important component in preventing and treating coagulopathy in trauma.
On average a unit contains a volume of 200-250 mL and is transfused rapidly.

1. Type A or AB thawed plasma is the current standard for transfusion during evacuation.

2. Thawed plasma only has a shelf life of 5 days and may not be available for the pre-hospital mission. Liquid plasma (never frozen) has a shelf life of 26 days. Check with issuing facility or blood supply unit for availability.

The recommended mission loads for tactical evacuation are based on operations tempo and determined by the theater or Joint Task Force surgeon. Specific missions may require additional blood products; units will refer to the Joint Blood Program Office.

1. Golden Hour Container (GHC) maximum capacity is 4 units RBC/FFP or 2 units of whole blood.

2. If whole blood or plasma is unavailable, evacuation personnel will fly with RBCs exclusively.

3. Product must finish infusion within 4 hours of spiking the bag. If not complete, then it needs to be stopped and the remainder of the product discarded. If the initiation of transfusion (spiking of the bag) is delayed, then the blood may be returned to storage if it hasn’t exceeded appropriate transport temperature, which is a max of 10C. The only way this can be determined is with use of a Safe-T-Vue sticker on the actual blood bag.

4. Unused blood products (i.e. WB and freeze dried plasma [FDP]) furnished by forward U.S. or Coalition Forces will not be used by evacuation personnel. Recommend products be left with forward forces. Blood products (WB and FDP) spiked by forward forces and transfusing at time of pick up will be continued during evacuation.

Pediatric patients:

1. Emergency transfusion of pediatric patients relies on clinical assessment rather than specific vital signs, since normal heart rate and blood pressure are age-dependent.

2. Clinical signs of shock are the same as in adults (cool, pale, weak or absent radial pulse, delayed capillary refill, decreased mental status).

3. Pediatric fluid resuscitation related to trauma begins with 10 mL/kg of first blood product, then repeat as needed based on response.
Receiving Blood Components from an Issuing Facility (U.S. and Coalition)

U.S. issuing facility personnel from the Blood Support Detachment (BSD), MTF (Role 2/3) or lab will:

1. If requested and available, thaw frozen plasma IAW local procedures and label products (A or AB) with 5 day expiration date.
2. Ensure GHC is properly charged and removed from freezer 25-30 minutes prior to loading blood products.
3. Ensure all blood products issued have a Safe-T-VUE (NSN 6515-08-T00-3056) attached and activated for temperature monitoring. *Ensure thawed plasma is at refrigerated temperature (1-6°C) before placing Safe-T-VUE on unit.*
4. Evacuation personnel will follow Safe-T-VUE procedures in Appendix E when required.
5. Document in TMDS the issuance of blood products to an evacuation team (e.g., DUSTOFF; Pararescue; Tactical Critical Care Evacuation Team).
6. Complete appropriate sections of the SF 518 Blood or Blood Component Transfusion Record for issuing blood products; place inside GHC pocket or attach form to each unit of blood product issued.
7. Verify the blood information on the SF 518 against the blood product label with receiving evacuation unit personnel.

Non-U.S. Issuing Facility: When U.S. blood products are to be issued from a Coalition facility, contact the Joint Blood Program Office to coordinate issuing requirements and documentation of units received.

Receiving unit (Evacuation Unit) personnel will:

1. Prior to sealing GHC, ensure each blood product loaded into the GHC has an activated Safe-T-VUE attached and an SF 518 Form.
2. Accept blood products into receiving unit’s TMDS inventory. *If receiving unit is unable to access TMDS, the issuing facility will access account and receive the products under the receiving unit’s TMDS inventory.*

Storage, Transportation and Monitoring of Blood Products

1. All blood and blood components must be maintained in a controlled environment and stored under appropriate conditions.
2. Blood products carried outside a BSD/MTF/Lab will only be transported in an approved storage container (e.g., Golden Hour Container NSN 6530-01-505-5301; OCP/5306; Desert) for a maximum of 48 hours.
3. Units will monitor containers and document status (e.g., dry/no leaking noted) at a minimum of every 24 hours.
4. Once loaded and sealed, container will remain closed and intact at all times until blood product is required for patient care.
5. Notify the issuing facility (BSD/MTF/Lab) as soon as possible when blood products have been used.

6. GHC is only approved for 48 hours use; prior to expiration end users will contact issuing facility (BSD/MTF/LAB) to coordinate the return and exchange of a container and blood products per mission requirements.

**WARNINGS:**
- At no time will container or its contents (blood products) be placed in a refrigerator or other cooling device outside a blood bank.
- Blood products will not be used if container is leaking; or the temperature indicator (Safe-T-VUE) on the blood product is out of standard (refer to Appendix E).
- Notify the issuing facility and return container and products for replacement.

**Individual and Unit Training Requirements**

1. At a minimum, medical personnel who participate in the administration of blood products during evacuation will be trained in the following topics:
   
   a. Indications for transfusion (Appendix C)
   
   b. Transfusion procedures (Appendix D);
   
   c. Documentation (Appendix G)
   
   d. PEARLS for transfusion (Appendix F)
   
   e. Submission of a patient safety report when required

2. At a minimum, non-medical personnel who assist will be trained in the following:
   
   a. Transfusion procedures
   
   b. Equipment/supplies
   
   c. Documentation requirements for the SF 518

3. Units who implement this CPG will train appropriate personnel on the following:
   
   a. Emergency procedures for in-flight complications.
   
   b. Storage container/blood product exchange requirements.

**Essential Items Required for Implementing a Vampire Program**

1. Approved blood component transport container.
   
   a. Recommend between 4 and 6 each GHCs for a Vampire Program (NSN 6530-01-505-5301 (OCP)/5306(Desert)).
b. Hemacool (NSN 4110-01-506-0895) or other freezer with temp check to ensure a temperature ≤ to (-) 18°C to support reconditioning of GHC.

2. Safe-T-VUE (NSN 6515-08-T00-3056) for temperature monitoring. (Refer to Appendix E.)

3. Establish Theater Medical Data Store (TMDS) accounts for an issuing facility (BSD/MTF (Role2/3) and LAB); and receiving unit (evacuation unit)

Warming Devices for Blood Transfusion

Use of infusion warming devices is highly recommended. These will be FDA approved for the actual use in transfusion of blood products (examples of devices include: Belmont® Buddy-lite™, EnFlow® or Thermal Angel).

**WARNING**: Warming devices will have safety mechanisms built in that prevents the output temperature from exceeding 42°C. Unit personnel will be familiar with safety mechanisms for the device used.

Tranexamic Acid (TXA)

Patients receiving blood transfusion within three hours of injury should also receive TXA. Refer to the TCCC guidelines for administration of TXA.

Record Keeping and Documentation Requirements

1. Transfusions will be documented into TMDS by evacuation personnel, or, by issuing facility.


3. Complete SF 518 documentation and turn over at the destination MTF for placement in the patient’s medical record.

References


APPENDIX C: CLINICAL INDICATIONS OF HEMORRHAGIC SHOCK

<table>
<thead>
<tr>
<th>Clinical Evidence Hemorrhagic Shock Is Present</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>H</strong> Hypotension</td>
</tr>
<tr>
<td><strong>T</strong> Tachycardia</td>
</tr>
<tr>
<td><strong>R</strong> Respirations</td>
</tr>
<tr>
<td><strong>P</strong> Pulse (poor character)</td>
</tr>
<tr>
<td><strong>M</strong> Mental status</td>
</tr>
<tr>
<td><strong>S</strong> Skin color</td>
</tr>
<tr>
<td><strong>C</strong> Continued bleeding</td>
</tr>
</tbody>
</table>
**APPENDIX D: TRANSFUSION PROCEDURES**

**TRANSFUSION PROCEDURES**

**MAINTAIN UNIVERSAL PRECAUTIONS (Gloves & Eye Protection)**

**STEP 1: ESSENTIAL BLOOD ADMINISTRATION ITEMS**

| 1. "Y" Type Filtered Blood Administration Set (UNDER NO circumstances should non-filtered tubing be used) | 4. Blood Pressure Cuff/Monitor |
| 2. Blood Product to Transfuse (Universal Donor is approved for Prehospital) | 5. Blood Warmer Device |
| 3. 0.9% NS (Dedicated Line Only for Blood Products) | 6. Pressure bag (if available) |

**STEP 2: PRE-TRANSFUSION TASK**

Two Person Verification Process

Verify Blood Label. Complete SF 518 for 5 items below or transcribe items from Blood Label onto blank SF 518:

1. Unit#;
2. Type of Product;
3. Donor ABO/Rh (Must be O for RBCs; and A or AB for Plasma);
4. Expiration Date; and
5. Temperature Indicator (RED = NOT ACCEPTABLE)

1. CLOSE all 3 clamps on Y tubing
2. NOTE: When using blood/fluid warming device, attach line to fluid warmer cartridge and fluid warmer extension line.
   a. Ensure warming device is functioning IAW manufacturers guidelines
3. Insert 1 spike into NS bag and hang; OPEN clamp and prime only the "Y" section; CLOSE clamp
4. Insert 2nd spike into blood product and hang; OPEN clamp and run the length of the tubing
5. Attach line to IV or 10 site **Ensure good flow through IV/10 before initiating transfusion**
6. Ensure all clamps are CLOSED
7. Note/document pre-transfusion vitals - at a minimum BP and HR
8. Medical person will visually inspect blood product if possible for gas, discoloration, clots, foreign objects, or, sediment, and ensure no cracking of the plastic bag that has led to leaking.
   a. Visually inspect the Temperature Indicator (RED = NOT ACCEPTABLE)
9. Non-Medical person can assist with documentation on the SF 518 for Pre and Post transfusion in formation

**STEP 3: TRANSFUSION TASK**

1. OPEN main line clamp for blood product to begin infusion.
   a. ENSURE CLAMP to NS REMAINS CLOSED.
   b. UNDER NO CIRCUMSTANCES will other medications or IV fluid (including 3%NS) be introduced through transfusion line. This will cause hemolysis/clotting of blood products.
2. **Blood products must be transfused within 4 hours of removal from a storage container** - if not, the product(s) will be returned to issuing facility or delivered with patient to MTF to be discarded.
3. If using pressure infuser set pressure to 300 mmHg.
5. When blood product has been infused, CLAMP blood product line and OPEN NS line to deliver residual blood product.
6. If 2nd Unit required - CLOSE NS clamp.
7. Spike 2nd Unit - OPEN blood product and main line clamps to begin 2nd infusion.
8. Monitor closely and continue VS assessment.
9. VS goal: SBP >100mmHg; and/or Pulse <100; MAP 70-80 mmHg

**STEP 4: DOCUMENTATION TASK**

1. Pre-transfusion Data
   a. Unit Number
   b. Type of Blood Product (RBC/Plasma)
   c. Donor ABO/Rh
   d. Expiration Date
   e. Vital Signs (HR and B/P)
2. Post Transfusion Data
   a. Vital Signs
   b. Date/Time started/completed
   c. Note if interrupted and reason for interruption
   d. Patient Identification (as much as possible)
## APPENDIX E: SAFE-T-VUE TEMPERATURE INDICATOR

### SAFE-T-VUE TEMPERATURE INDICATOR

#### ISSUING FACILITY INSTRUCTIONS

1. Safe-T-VUE® 10 is a temperature sensitive indicator that easily adheres directly to blood bags during transport and changes color from WHITE to RED when the 10°C indication temperature has been reached or exceeded.
   a. Safe-T-VUE is non-reversible and indicates that a high temperature condition existed, even if temperature returns to a lower level. As long as indicator remains WHITE, blood may be stored for future use.
2. Prepare the Safe-T-VUE temperature indicator by refrigerating for a minimum of 24 hours at 1-6°C.
3. Remove the blood product and one Safe-T-VUE indicator from the refrigerator at the same time and place on a clean dry surface.

**NOTE:** Remove excess moisture from the blood product bag by using a dry wipe/paper towel on the surface where the Safe-T-VUE is to be applied.

**NOTE:** Use of cold pack on the surface below the blood product will help to maintain temperature

4. Hold Safe-T-VUE against the blood product with finger tips. Peel off the “REMOVE” label to expose the adhesive.

**NOTE:** Be careful to only handle around the edge of the indicator to expose RED DOT and WHITE DOT.

5. Attach Safe-T-VUE directly to the lower third of the blood product bag where there is a large volume of product.

**CAUTION:** Ensure thawed plasma is at refrigerated temperature (1-6°C) before placing Safe-T-VUE on unit.

**NOTE:** Be certain the Safe-T-VUE indicator is in complete contact with the blood component bag being monitored. No air pockets should be under the indicator (e.g., fold in the bag; over any labels; or any other obstruction).

6. Fold WHITE DOT onto the RED DOT and press firmly together to activate.

**CAUTION:** Be careful to ONLY press on the GREEN color-coded end to activate properly.

**CAUTION:** It is important to place pressure on the outer edge of the WHITE DOT, and not the center, when pressing onto the RED DOT to prevent false activation.

7. Issuing facility will complete documentation on SF 518 for each blood product unit, place inside GHC pocket and secure container.

8. Receiving personnel will understand color change temperature indication:
   a. When WHITE DOT turns solid RED, temperature has reached ≥10°C
      • Return blood product to issuing facility (BSD/MTF/LAB)
   b. Appearance of SMALL RED DOTS is an indication blood product requires cooling or immediate refrigeration.
      • Return product to issuing facility (BSD/MTF/LAB) for appropriate cooling/refrigeration
   c. WHITE DOT – product is acceptable for transfusion

![Temperature Indicator Diagram]
# PEARLS FOR TRANSFUSIONS

## PRE-TRANSFUSION PEARLS

1. Use of 2% Lidocaine (2-3ml) with 0.9% NS is permitted to flush any IO site prior to blood product transfusion.
2. Consider pain control measures to reduce tachycardia resulting from uncontrolled pain.
3. Once removed from storage container blood products will be transfused in under 4 hours.
4. ONLY USE “Y” filtered blood administration sets.
5. If directly involved in patient care, 1st Verifier (Medical Person) can direct a non-medical person to be the 2nd Verifier and record data on the SF518.
6. **If using enFlow® fluid warmer – add IV extension tubing**
7. DO NOT use blood product if storage container is leaking or temperature indicator is RED.
8. DO NOT allow blood warmer to be placed directly on patients skin as this may cause burning.
9. If Thawed plasma is available it should be given prior to RBC; normal ratio is 1:1.

## DURING TRANSFUSION PEARLS

1. Transfusion infusion rates can be titrated to slower rates if VS parameters move to appropriate levels (SBP>100; HR<100; MAP 70-80).
2. Special attention should be paid to non-compressible injuries (chest; abdominal; and pelvis) so as to NOT raise the SBP over 90mmHg.
3. Once transfusion is initiated, decrease all other fluids to KVO rate.
4. **In-flight emergencies:**
   a. Contact unit FS or tactical operation center for medical direction; or
   b. Divert to nearest MTF (Do not delay divert waiting on medical direction).
5. If transfusion is interrupted, record date/time and reason for interruption on SF518 if not able to resume within 5 min.
6. Under NO CIRCUMSTANCES will other medications or IV fluids (to include 3% NS) be introduced through transfusion line.
7. Blood output temperature from a warmer device WILL NOT EXCEED 42°C (107°F).

## DURING TRANSFUSION PEARLS

1. Suspected /confirmed transfusion reaction: STOP TRANSFUSION
2. Disconnect tubing from infusion site; flush IV site with NS
3. Keep IV Line OPEN with NS
4. Re-initiate transfusion only if it is deemed clinically essential
5. Document on SF 518 date/time and actions taken

## POLE TRANSFUSION PEARLS

1. After 1st transfusion, re-evaluate casualty and initiate 2nd unit ONLY if criteria is still met (Appendix A).
2. If 1st Unit is initiated based on “Stand-Alone” injury (Double/Triple/Quadruple Amputation); subsequent units will be based on VS parameters.
3. Complete documentation on SF518.
4. Consider Tranexamic Acid (TXA) – follow TCCC Guidelines for Administration.

## PATIENT HAND-OFF (COMMUNICATION)

1. Provide receiving MTF with completed SF 518s for patients record.
2. Report any adverse events; transfusion reactions; and actions taken enroute.
3. Report interrupted transfusions and provide explanation.
4. Report O POS blood given to female patients between the age of 10-50.
APPENDIX G: REQUIRED DOCUMENTATION FOR TRANSFUSION

- **SF 518 Blood or Blood Component Transfusion Record ASBP**
- **151 Whole Blood Transfusion Checklist**
- **147 Eldon Card ABO/Rh Typing Record**
- **145 Infectious Disease Testing for Blood Donation**
- **148 Pre-screen Whole Blood Sample Shipping Manifest**
- **150A Emergency Release Letter of Understanding (tested)**
- **150B Emergency Release Letter of Understanding (untested)**
- **572 Emergency Whole Blood Donation Record**
- **AABB Medications Deferral List**
- **DHQ Medication Deferral List**
APPENDIX H: ADDITIONAL INFORMATION REGARDING OFF-LABEL USES IN CPGS

PURPOSE

The purpose of this Appendix is to ensure an understanding of DoD policy and practice regarding inclusion in CPGs of “off-label” uses of U.S. Food and Drug Administration (FDA)–approved products. This applies to off-label uses with patients who are armed forces members.

BACKGROUND

Unapproved (i.e. “off-label”) uses of FDA-approved products are extremely common in American medicine and are usually not subject to any special regulations. However, under Federal law, in some circumstances, unapproved uses of approved drugs are subject to FDA regulations governing “investigational new drugs.” These circumstances include such uses as part of clinical trials, and in the military context, command required, unapproved uses. Some command requested unapproved uses may also be subject to special regulations.

ADDITIONAL INFORMATION REGARDING OFF-LABEL USES IN CPGS

The inclusion in CPGs of off-label uses is not a clinical trial, nor is it a command request or requirement. Further, it does not imply that the Military Health System requires that use by DoD health care practitioners or considers it to be the “standard of care.” Rather, the inclusion in CPGs of off-label uses is to inform the clinical judgment of the responsible health care practitioner by providing information regarding potential risks and benefits of treatment alternatives. The decision is for the clinical judgment of the responsible health care practitioner within the practitioner-patient relationship.

ADDITIONAL PROCEDURES

Balanced Discussion

Consistent with this purpose, CPG discussions of off-label uses specifically state that they are uses not approved by the FDA. Further, such discussions are balanced in the presentation of appropriate clinical study data, including any such data that suggest caution in the use of the product and specifically including any FDA-issued warnings.

Quality Assurance Monitoring

With respect to such off-label uses, DoD procedure is to maintain a regular system of quality assurance monitoring of outcomes and known potential adverse events. For this reason, the importance of accurate clinical records is underscored.

Information to Patients

Good clinical practice includes the provision of appropriate information to patients. Each CPG discussing an unusual off-label use will address the issue of information to patients. When practicable, consideration will be given to including in an appendix an appropriate information sheet for distribution to patients, whether before or after use of the product. Information to patients should address in plain language: a) that the use is not approved by the FDA; b) the reasons why a DoD health care practitioner would decide to use the product for this purpose; and c) the potential risks associated with such use.