Pre-hospital transfusion of plasma in hemorrhaging trauma patients independently improves hemostatic competence and acidosis

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Abstract

Background: The early use of blood products has been associated with improved patient outcomes following severe hemorrhage or traumatic injury. We aimed to investigate the influence of pre-hospital blood products (i.e. plasma and/or RBCs) on admission hemostatic properties and patient outcomes. We hypothesized that pre-hospital plasma would improve hemostatic function as evaluated by rapid thrombelastography (rTEG).

Methods: We conducted a prospective observational study recruiting 257 trauma patients admitted to a Level I trauma center having received either blood products pre-hospital or in-hospital within 6 hours of admission. Clinical data on patient demographics, blood biochemistry, injury severity score and mortality were collected. Admission rTEG was conducted to characterize the coagulation profile and hemostatic function.

Results: 75 patients received pre-hospital plasma and/or RBCs (PH group; nearly half received both RBCs and plasma) whereas 182 patients only received in-hospital blood products (RBCs, Plasma and Platelets) within 6 hours of admission (IH group). PH patients had lower Glasgow coma scale (GCS) scores, more penetrating injuries, lower systolic blood pressures, lower hemoglobin levels, lower platelet counts and greater acidosis upon ED admission than the IH group (all \( p < 0.05 \)). Despite differences in type of injury and admission vitals indicating that the PH group had more signs of bleeding than the IH group, there were no significant differences in in-hospital mortality (PH 26.7% vs. IH 20.9% \( p = 0.31 \)). When comparing rTEG variables between PH patients transfused with 0, 1 or 2 units of plasma, more pre-hospital plasma transfusion was tendency towards improved rTEG variables. When adjusting for pre-hospital RBC, pre-hospital plasma was associated with significantly higher rTEG MA (\( p = 0.012 \)) at hospital admission.

Discussion: After adjusting for pre-hospital RBCs, pre-hospital plasma transfusion was independently associated with increased rTEG MA, as well as arrival indices of shock and hemodynamic instability. Besides more severe injury and worse clinical presentation, the group that received pre-hospital transfusion had early and late mortality similar to patients not transfused pre-hospital.

Conclusions: These data suggest that early administration of plasma can provide significant hemostatic and potential survival benefit to severely hemorrhaging trauma patients.

Keywords: Pre-hospital, Plasma, RBCs, Trauma, Transfusion, Thrombelastography, Coagulopathy
Background

Death from injury has increased by 20% over the last decade and accounts for more deaths than malaria, tuberculosis and HIV combined [1]. Hemorrhage requiring massive transfusion secondary to trauma remains a major cause of potentially preventable deaths, and development of trauma induced coagulopathy (TIC) further increases the mortality rates of hemorrhaging patients substantially [2]. During the last decade a number of strategies and protocols have been developed and adopted to improve the management and delivery of care to trauma patients with severe hemorrhage. These strategies include: (i) damage control surgery, the use of abbreviated surgery to remedy hemorrhage and contamination with definitive surgery delayed until physiology has been restored [3]; (ii) limited use of crystalloid fluids, which can worsen coagulopathy through dilution of clotting factors and platelets [4]; (iii) hypotensive resuscitation, whereby normalization of blood pressure is delayed until hemorrhage has been surgically controlled [5]; and (iv) balanced blood product use, whereby packed red blood cells (RBCs), plasma and platelets are transfused in ratios of, or near to, 1:1:1 mimicking whole blood, which is associated with improved outcomes and may help alleviate TIC [6, 7]. Taken together, these are known as damage control resuscitation (DCR) [8]. As DCR developed, it became clear that early, aggressive resuscitation with blood products could lead to early correction of acidosis, coagulopathy, and hypothermia [9–11].

Rawdan and colleagues reported that moving thawed plasma from the blood bank to the emergency department enabling early balanced blood product transfusions, was associated with a reduction in overall blood product use and a 60% decrease in odds of 30-day mortality [12]. Based on this rapidly evolving experience, the Texas Trauma Institute at Memorial Hermann Hospital (Houston, TX) introduced the availability of two units of O RhD negative RBCs and two units of thawed AB plasma in appropriate temperature controlled containers on four Life Flight (LF) helicopters [13]. Recently, we reported that this novel practice of early hemostatic resuscitation with blood products was associated with improved hemorrhage-resuscitation and patient outcomes (i.e. improved survival) when compared to trauma patients only receiving crystalloids en route to the hospital [13].

The objective of the present observational study was to investigate the association between pre-hospital administered RBCs and plasma and hemostatic function as evaluated by whole blood thrombelastography (TEG) on arrival at the trauma center as compared to that of patients not receiving pre-hospital blood transfusions. We hypothesized that patients receiving pre-hospital blood products, specifically pre-hospital plasma, would display improved hemostatic function as evaluated by TEG.

Methods

Setting and patients

The Texas Trauma Institute at Memorial Hermann Hospital is an American College of Surgeons verified Level I trauma center that is the primary teaching hospital for the University of Texas Health Science Center and operates a hospital-based helicopter program, Life Flight (LF). The helicopters carry two units of type O RBCs and two units of AB plasma for emergent use following established guidelines. This prospective observational cohort study included 257 trauma patients admitted directly from the scene of the injury to the highest level of activation at Memorial Hermann Hospital Texas Medical Center (MHH-TMC) between October 2012 and November 2013. The University of Texas Health Science Center at Houston institutional review board (IRB) approved this study (IRB# HSC-GEN-12-0059).

Patient selection

All adult trauma patients (≥16 years) who met criteria for full trauma team activation and received blood before arrival at ED (i.e. pre-hospital, PH group) or after hospital admittance within 6 h of ED arrival (i.e. institutional, IH group) were included in this study.

The criteria for exclusion were: pregnancy, prisoner, had a burned body surface area >20% or enrolled in another study. Patients from whom we could not obtain a blood draw were excluded from analysis. Informed consent was obtained from the patient or a legally authorized representative within 72 h of admission. If the patients were discharged or died within 24 h, a waiver of consent was granted. In cases which consent could not be obtained, the patient was excluded from the study, and their blood samples were destroyed. Patient information on baseline demographics, clinical, physiologic, and laboratory tests were extracted from patient records and the institutional trauma registry.

Sample collection

Blood samples were collected immediately upon hospital admission on all adult trauma patients (≥16 years) who met criteria for full trauma team activation i.e., PH patients had samples drawn after PH blood transfusion whereas IH patients had samples drawn before any blood transfusion. In all cases, blood was drawn into citrated tubes for conducting the rapid-TEG assay.

Rapid Thrombelastography (rTEG)

All rTEG specimens were run on a thrombelastograph 5000 (Haemoscope Corporation, Niles, IL). Admission rTEGs were performed using the accepted citrate reversal method. Specimens were collected in a small (3 mL) citrated tube, transported to the Emergency Department Stat Laboratory. The citrate was immediately reversed.
by adding calcium chloride according to the recommenda-
tions of the manufacturer within the rTEG package
insert. After this, TEG was performed within 5–10 min
of blood sampling, using tissue factor and kaolin as acti-
vators. Staff laboratory technicians in the Memorial Her-
mann Hospital Emergency Department performed all
the rTEG assays during the defined study period. These
same technicians performed all the quality controls on
the TEG analyzers, doing so every 8 h. Quality control
was performed per the package insert from the
manufacturer.

The rTEG generates several values that describe the
hemostatic process: The first value generated is the ac-
tivated clotting time (ACT), which is the time in seconds
between initiation of the test and the initial fibrin for-
mation (normal range, 86–118 s). Similar to the ACT, the
r-value (also known as the reaction time (R-time), 0–1
min) expresses the time between the start of the assay
and beginning of clot formation. The K-time (normal
range, 1–2 min) is the time needed to reach 20-mm clot
strength and the alpha (α) angle (normal range, 66–82°)
determines the slope of the tracing that represents the
rate of clot formation. The maximal amplitude (MA;
normal range, 54–72 mm) is the greatest amplitude of
the tracing and the G-value (normal range, 5,300–12,000
dynes/cm²) is a global measure of absolute clot strength
(both enzymatic and platelet contributions) and is de-
creased in hypocoagulable states. LY30 (normal range,
0.0–7.5%) is the percent amplitude reduction at 30 min
after MA, illustrating the amount of fibrinolysis.

Statistics
Statistical analysis was performed using STATA (IC ver-
sion 12.1, College Station, TX) and SPSS 22 (IBM Cor-
nporation, New York, US). Descriptive data are presented
as medians with inter quartile ranges (IQR) or as %
 proporitions).

Non-parametric statistical tests (Mann-Whitney U test
and Kruskal-Wallis) and Chi-square tests were used to
evaluate group differences (PH vs. IH groups), as appro-
priate. P-values <0.05 were considered significant.

To investigate the association of pre-hospital RBCs
and plasma on admission for rTEG MA, a parameter
with significant predictive value for transfusion require-
ments and mortality [14–16], we conducted a linear re-
gression analysis. Univariate and multivariate linear
regression models were constructed for each group (PH,
IH) and combined including both groups. Multivariate
linear regression models were adjusted for admission
 vital differences between the PH and IH groups (systolic
blood pressure (SBP), hemoglobin, platelet count and
pH) as well as adjusting for mode of transport (i.e. LF or
ambulance) and amount of PH-RBC and PH-Plasma in
the model including both groups (PH and IH).

Furthermore we compared rTEG parameters in PH pa-
tients receiving plasma based on the amount adminis-
tered (0, 1 and 2 units plasma) by ANOVA and Tukey
post-hoc test.

Results
Patient characteristics
A total of 257 patients were included in the study, 75 (29%)
patients received pre-hospital (PH) blood, whereas 182
(71%) patients only received blood in-hospital (IH) within
6 h of hospital admission. Eighty-six patients were trans-
ported by ground ambulance and 171 patients by LF, of
which 44% received a PH blood transfusion (Table 1).
The median age of the study cohort was 37 years, 81% were
male, with a median injury severity score (ISS) of 26
(Table 1). Patients in the PH group demonstrated a higher
incidence of penetrating injuries (45 vs. 25%, p = 0.002),
field intubations (39 vs. 21%, p = 0.005), positive FAST
exams and presented with lower GCS scores compared to
patients in the IH group (Table 1). Additionally, patients re-
ceiving PH blood displayed significantly lower systolic
blood pressure (SBP), hemoglobin, platelet count and more
acidosis than IH, suggesting a more pronounced degree of
hemorrhage and/or shock in the PH group. Lastly, patients
receiving PH blood received more units of blood products
within 6 and 24 h after admission (Table 1). Despite differ-
ences in type of injury and admission vitals indicating more
severe injury and bleeding in the PH group, there were no
 significant differences in mortality (Table 1).

Pre-hospital Plasma is associated with improved
hemostasis
Despite the variations between the two groups, no signif-
 icant differences were found in the median rTEG values,
respectively for the PH vs. IH, for ACT (121 vs. 121), R-
time (0.8 vs. 0.8), K-time (1.65 vs. 1.45), Angle (70 vs. 73)
and Ly30 (1 vs. 1.4), upon arrival at the hospital. However,
at hospital admission PH displayed significant lower me-
dian rTEG MA (62 vs. 64, P = 0.020) and G-value (8.1 vs.
8.69, P = 0.009) compared to IH patients. Table 2 shows
the distribution of pre-hospital blood products transfused.
Nearly half of the PH group received 1 unit of RBCs and
plasma each (Table 2). When comparing rTEG values
ACT, R-time, K-time, α-Angle, MA, G and Ly30 in PH pa-
tients receiving 0, 1 and 2 units of plasma pre-hospital un-
adjusted for PH RBC, patients receiving 2 units of plasma
pre-hospital had the highest mean values of rTEG α-Angle
(70.57 vs. 67.51 vs. 72.00° for plasma = 0, 1 and 2 units,
respectively), rTEG MA (58.73 vs. 57.30 vs. 62.55 mm for
plasma = 0, 1 and 2 units, respectively) and rTEG G (5.93
vs. 7.36 vs. 8.41 dyn/cm² for plasma = 0, 1 and 2 units,
respectively). Furthermore, patients receiving 2 units of
plasma pre-hospital had lower average values of K-time
(1.63 vs. 2.07 vs. 1.59 min for plasma = 0, 1 and 2 units,
respectively) and R-time (2.07 vs. 1.07 vs. 0.72 min for plasma = 0, 1 and 2 respectively). There was a significant difference in R-time between patients receiving no pre-hospital plasma compared to patients receiving 2 units of plasma ($P = 0.023$).

**Predictors of hemostatic competence**

In the PH group, the unadjusted univariate linear regression demonstrated that pre-hospital plasma was

### Table 1 Patient demographics, injury severity, admission vitals, transfusion and outcome in 257 adult trauma patients

|                     | Units | Overall (n = 257) | PH (n = 75) | IH (n = 182) | P-value |
|---------------------|-------|------------------|-------------|--------------|---------|
| Age                 | Years | 37 (25, 53)      | 34 (23, 53) | 39 (26, 53)  | 0.259   |
| Male                | n (%) | 208 (80.9%)      | 62 (82.7%)  | 146 (80.2%)  | 0.650   |
| Injury Scores and severity |       |                 |             |              |         |
| ISS Score           |       | 26 (17, 34)      | 29 (17, 41) | 26 (17, 34)  | 0.106   |
| GCS Score           |       | 10 (3, 15)       | 3 (3, 15)   | 12 (3, 15)   | 0.022   |
| Positive FAST %     |       | 54 (31.4%)       | 26 (48.1%)  | 28 (23.7%)   | 0.001   |
| Penetrating Injury %|       | 80 (31.1%)       | 34 (45.3%)  | 46 (25.3%)   | 0.002   |
| Intubated in the field |     | 63 (27.2%)       | 29 (39.2%)  | 34 (21.5%)   | 0.005   |
| Admission Vitals    |       |                 |             |              |         |
| SBP mmHg            |       | 98 (80, 123)     | 90 (77, 113)| 100 (80, 125)| 0.044   |
| DBP mmHg            |       | 60 (48, 74)      | 59 (50, 69) | 60 (48, 76)  | 0.299   |
| Heart rate Bpm      |       | 108 (88, 130)    | 111 (90, 133)| 108 (85, 130)| 0.425   |
| Hemoglobin g/dL     |       | 12.9 (11.3, 14.2)| 12.4 (11.3, 13.4)| 13.1 (11.35, 14.3)| 0.011   |
| Platelet count x10^9/μL | 215 (163, 250) | 193 (152, 223)  | 225 (174, 257) | 0.001   |
| pH                  |       | 7.26 (7.15, 7.32) | 7.21 (7.06, 7.32)| 7.27 (7.18, 7.33) | 0.002   |
| Base Excess mEq/L   |       | −5 (−10, −2)     | −6 (−10, −3) | −4 (−10, −1) | 0.073   |

**Mode of Transport**

|                     | LF %  | 171 (66.8%) | 75 (100%) | 96 (52.7%) | <0.001 |

**Transfusion: Prehospital**

| RBC Units          | 0 (0, 0) | 1 (1, 2) | 0 (0, 0) | NA |
| Plasma Units       | 0 (0, 0) | 1 (1, 2) | 0 (0, 0) | NA |

**Transfusion: Admission - 6 h total transfusions**

| RBC Units          | 4 (2, 10) | 8 (4, 14) | 3 (1, 8) | <0.001 |
| Plasma Units       | 4 (2, 8)  | 6 (3, 12) | 3 (2, 7) | <0.001 |
| Platelets Packs    | 0 (0, 6)  | 6 (0, 12) | 0 (0, 6) | <0.001 |

**Transfusion: Admission - 24 h total transfusions**

| RBC Units          | 5 (2, 12) | 10 (4, 15) | 4 (2, 9) | <0.001 |
| Plasma Units       | 4 (2, 9)  | 7 (3, 14) | 4 (2, 8) | <0.001 |
| Platelets Packs    | 6 (0, 12) | 6 (0, 18) | 0 (0, 6) | <0.001 |

**Outcomes**

| 6-h mortality %    | 25 (9.7%) | 10 (13.3%) | 15 (8.2%) | 0.210 |
| 24-h mortality %   | 31 (12%)  | 12 (16%)   | 19 (10.4%)| 0.213 |
| In-hospital mortality % | 58 (22.6%) | 20 (26.7%) | 38 (20.9%)| 0.313 |

Patients met criteria for full trauma team activation and either received blood before arrival at ED (PH) or in the first 6 h after hospital admission (IH). Medians (IQR) or n (%) are reported. All the bold numbers reflect data that is significant.

### Table 2 Distribution of patients receiving pre-hospital blood products

| Plasma | 0 U | 1 U | 2 U | Total RBC |
|--------|-----|-----|-----|-----------|
| 0 U    | 0 (0%) | 10 (13%) | 3 (4%) | 13 (17%) |
| RBC    | 1 U | 2 (3%) | 30 (40%) | 7 (9%) | 39 (52%) |
| 2 U | 1 (1%) | 2 (3%) | 20 (27%) | 23 (31%) |
| Total Plasma | 3 (4%) | 42 (56%) | 30 (40%) | 75 (100%) |
associated with significantly higher rTEG MA values (one unit increase in PH plasma was associated with (β coefficient) 13.95 mm, 95%CI [3.13, 24.77], p = 0.012) higher MA at hospital admission in a model only adjusting for PH RBC. The interaction between PH RBC and plasma did not appear to have a significant effect on admission rTEG MA values (data not shown). When adjusting for pH, hemoglobin, platelet count, SBP, and in PH patients PH-RBC and PH-Plasma, higher pH and higher platelet count were independent predictors of rTEG MA in IH patients and higher pH, higher platelet count and PH-Plasma were associated with higher rTEG MA in PH patients (all p < 0.05, data not shown).

When conducting the analysis non-stratified (including both PH and IH groups in the same analysis), and instead adjusting for transportation (by LF or ambulance) and admission vital differences, and PH transfusion with Plasma and RBC, PH plasma transfusions were associated independently with higher rTEG MA values together with higher pH, higher SBP and higher platelet count (Table 3).

Discussion

This study found that after adjusting for pre-hospital RBCs, pre-hospital plasma transfusion was independently associated with increased rTEG MA, as well as arrival indices of shock and hemodynamic instability. Acidosis and low platelet count were found to impede hemostatic potential evaluated by rTEG. Besides more severe injury and worse clinical presentation, the group that received pre-hospital transfusion had early and late mortality similar to patients not transfused pre-hospital.

Reduced clot strength, as evaluated by TEG, has repeatedly been associated with increased bleeding, transfusion requirements and mortality in severely injured trauma patients [16, 17]. In the present study, plasma transfusion PH was independently associated with higher rTEG MA values, also after adjusting for pre-hospital RBCs. Furthermore, patients in the PH group receiving the highest amount of pre-hospital plasma transfusions differed in their hemostatic competence as evaluated by rTEG when compared to the patients in the PH group only receiving one or zero pre-hospital plasma. Given that the most severely injured and bleeding patients are expected to receive the highest amount of plasma pre-hospital, this suggests that pre-hospital plasma is more beneficial than pre-hospital RBCs for improving hemostasis in the acute/pre-hospital setting, in accordance with the coagulation factor and fibrinogen content in plasma compared to RBC.

Furthermore, PH patients were significantly different from IH patients. PH patients presented with more penetrating injuries, more positive FAST exams, lower systolic blood pressures, lower hemoglobin, lower platelet count, lower pH upon arrival to the trauma bay and lower rTEG MA and G, despite receiving pre-hospital RBC and/or plasma transfusions during transport. However, 52.7% of the IH patients were transported by LF where blood products are available; indicating that the prehospital blood transfusion by LF was not deemed necessary for these patients as they were not transfused. Furthermore, IH patients only received three RBCs and three plasma units within six hours of hospital admission, which suggests only moderate hemorrhage compared to eight RBCs and six plasma units in the PH group. Due to these differences, we stratified by group as well as conducting adjusted regression analyses. Regardless of group, however, acidosis and low platelet count were the strongest predictors of reduced hemostatic competence evaluated by rTEG MA values. This indicates that future analyses of rTEG should consider adjusting for pH and platelet count to make appropriate comparisons.

In addition, the acid buffer capacity of plasma is known to be 25–50 times better than crystalloids solutions [18], and despite the divergence between the two groups, the data shows no significant difference in the 6-h, 24-h, and the overall mortality. This indicates that prehospital plasma transfusion may lead to early correction of acidosis thereby mitigating coagulopathy, which

Table 3 Admission rapid thrombelastography (rTEG) MA-value evaluated by linear regression in 257 adult trauma patients

| Variable      | Units | Univariate β (95% CI) | P value | Adjusted β (95% CI) | P value |
|---------------|-------|-----------------------|---------|---------------------|---------|
| LF            | Yes   | −0.52 (−3.25, 2.21)   | 0.709   | 1.02 (−1.41, 3.46)  | 0.408   |
| pH            | mEq/l | 30.00 (21.53, 38.47)   | <0.001  | 23.55 (15.26, 31.8) | <0.001  |
| Hemoglobin    | g/dl  | 0.80 (0.19, 1.42)     | 0.011   | −0.12 (−0.68, 0.43) | 0.666   |
| Platelet counts | 10⁵/μL | 0.07 (0.05, 0.08)     | <0.001  | 0.06 (0.04, 0.08)  | <0.001  |
| SBP           | mmHg  | 0.07 (0.04, 0.11)     | <0.001  | 0.04 (0.00, 0.07)  | 0.047   |
| PH-RBC        | units | −1.89 (−3.86, 0.09)   | 0.061   | −1.82 (−4.82, 1.18) | 0.232   |
| PH-Plasma     | units | −0.85 (−2.68, 0.99)   | 0.364   | 2.88 (0.10, 5.66)   | 0.042   |

Regression coefficients (β) with 95% confidence intervals (95% CI) and p-values are displayed. P-values < 0.05 are shown in bold. Predicted change in rTEG MA (mm) associated with: LF (yes) and one unit higher pH, Hemoglobin, Platelet count, SBH, PH-RBC and PH-Plasma.
potentially can be translated into improved survival though this has to be investigated in future studies designed specifically to answer this question.

The study has several limitations. First, it should be noted that we could not obtain blood samples from “in-the-field”, prior to when the patient received pre-hospital transfusions. As such, we cannot definitively assess the effect of each transfusion and potential changes in the TEG values from “in-the-field” vs. hospital admission. Future studies are needed to evaluate these prehospital and pre-transfusion acute changes in clotting properties. Furthermore, another limitation is the relatively small sample size of 75 patients within the PH group and nearly half of these patients received one unit of RBCs and plasma pre-hospital. This emphasizes the need for future studies investigating the relative benefits of adding RBCs to plasma in a larger number of patients, preferably sampling blood for pre-hospital analysis.

Conclusion

In conclusion, after adjusting for pre-hospital RBCs, prehospital plasma transfusion was independently associated with higher rTEG MA. Acidosis (low pH levels) and low platelet counts were the strongest predictors of poor hemostasis evaluated by rTEG. Besides, more severe injury and worse clinical presentation, the PH group had early and late mortality similar to the IH group. Overall, these data imply that early administration of plasma may have beneficial effects on improving hemostasis, which may potentially translate into improved patient survival in future studies with larger sample sizes.

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None.

Authors’ contributions

All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

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