Comparing hemostatic resuscitation management of intraoperative massive bleeding with traumatic massive bleeding: a computer simulation

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Background: Appropriate blood component transfusion might differ between intraoperative massive bleeding and traumatic massive bleeding in the emergency department because trauma patients initially bleed undiluted blood and replacement typically lags behind blood loss. We compared these two blood loss scenarios, intraoperative and traumatic, using a computer simulation.

Methods: We modified the multi-compartment dynamic model developed by Hirshberg and implemented it using STELLA 9.0. In this model, blood pressure changes as blood volume fluctuates as bleeding rate and transcapillary refill rate are controlled by blood pressure. Using this simulation, we compared the intraoperative bleeding scenario with the traumatic bleeding scenario. In both scenarios, patients started to bleed at a rate of 50 ml/min. In the intraoperative bleeding scenario, fluid was administered to maintain isovolemic status; however, in the traumatic bleeding scenario, no fluid was supplied for up to 30 min and no blood was supplied for up to 50 min. Each unit of packed red blood cells (PRBC) was given when the hematocrit decreased to 27%, fresh frozen plasma (FFP) was transfused when plasma was diluted to 30%, and platelet concentrate (PC) was transfused when platelet count became 50,000/ml.

Results: In both scenarios, the appropriate ratio of PRBC:FFP was 1:0.47 before PC transfusion, and the ratio of PRBC:FFP:platelets was 1:0.35:0.39 after initiation of PC transfusion.

Conclusion: The ratio of transfused blood component did not differ between the intraoperative bleeding and traumatic bleeding scenarios.

Keywords: Blood coagulation disorder; Blood component transfusion; Computer simulation; Hemorrhage.
Table 1. Blood Component Information in the Transfusion Guidelines, Korea Center for Disease Control and Prevention, 2013

<table>
<thead>
<tr>
<th>Component</th>
<th>PRBC</th>
<th>FFP</th>
<th>Platelets</th>
</tr>
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<tbody>
<tr>
<td>Volume (ml)</td>
<td>243.12 ± 13.48</td>
<td>155.03 ± 12.11</td>
<td>50.85 ± 1.90</td>
</tr>
<tr>
<td>Hematocrit (%)</td>
<td>70.31 ± 4.16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Count (× 10^9/unit)</td>
<td></td>
<td></td>
<td>66.7 ± 11.7</td>
</tr>
</tbody>
</table>

Values are presented as mean ± SD. PRBC: packed red blood cells, FFP: fresh frozen plasma.
13.48 ml. Hematocrit was 70.31% on average. As a result, red blood cell volume was at an average of 170 ml. We assumed all PRBCs were derived from 400 ml of whole blood for convenience. In the manufacturing blood compartment, 56 ml of citrate-phosphate-dextrose with adenine (CPDA-1) solution was added. Plasma volume in PRBCs was 243 – 170 = 73 ml, which includes some amount of CPDA-1 solution. Total plasma volume in whole blood including CPDA-1 solution was 73 + 155 + 51 = 279 ml. Volume of CPDA-1 solution in PRBCs was 73 × 56 / 279 = 15 ml. Calculated real plasma volume contained in PRBCs was 73 – 15 = 58 ml, which excluded the volume of CPDA-1 solution.

The average volume of plasma derived from 400 ml whole blood was 155.03 ± 12.11 ml. Except for the volume of CPDA-1 solution, each fresh frozen plasma (FFP) was assumed to contain 124 ml of plasma. The average volume of platelet concentrate (PC) derived from 400 ml whole blood was 50.85 ± 1.90 ml. Except for the volume of CPDA-1 solution, each unit of PC contained 41 ml of plasma and 6.67 × 10^10 platelets on average, of which one third underwent splenic sequestration when transfused [16].

The reference scenario

We generated an intraoperative bleeding scenario and a trauma bleeding scenario. Both scenarios included a 70 kg person whose blood volume, hematocrit, fibrinogen concentration, and platelets of 4.9 L, 42%, 3 g/L, and 230,000/ml was assumed, respectively. In both scenarios, patients started to bleed at a rate of 50 ml/min.

The intraoperative bleeding scenario was based on a washout equation by designing the normovolemic status to be maintained by administering fluid. As bleeding continued, the blood pressure was assumed to be unchanged. If hematocrit decreased to 27%, one unit of PRBCs was given for 10 min. FFP was transfused for 10 min when plasma was diluted to 30% of the original plasma. Each unit of PC was transfused for 5 min when platelet count became 50,000/ml. FFP and PC starting time followed the practice guidelines for perioperative blood transfusion and adjuvant therapies [17].

In the traumatic bleeding scenario, the patient was set to bleed for 30 min without fluid supply until arrival at the emergency room (ER). The traumatic bleeding scenario was not only based on a washout equation, but also adopted a pressure-volume relationship concept. Using the pressure-volume equation, the amount of bleeding per minute reduced as blood pressure decreased due to a loss of blood. From arrival in the ER, crystalloid was administered via an intravenous line at the rate of 300 ml/min until the patient recovered to isovolemic status. After 50 min of trauma, blood components were prepared and each unit of PRBCs was given when hematocrit decreased to 27%. FFP was transfused when plasma was diluted to 30%, and PC was transfused when platelet count became 50,000/ml. The infusion of each blood component was same as in the intraoperative bleeding scenario.

RESULTS

In the intraoperative bleeding scenario (Fig. 2), bleeding during operation began at time 0 and PRBC administration started at 43 min when the bleeding volume was 0.44 times total blood volume (Table 2). In the traumatic bleeding scenario (Fig. 3), the patient lost 1,175 ml of blood volume and transcapillary refill volume was 449 ml until arrival in the ER when the systolic blood pressure was 84 mmHg. PRBC administration was started at 52 min when bleeding volume was 0.42 times total blood volume. In the intraoperative bleeding scenario, FFP administration started at 164 min when bleeding volume was 1.67 times the total blood volume. In the traumatic bleeding scenario, FFP administration started at 170 min when bleeding volume was 1.67 times the total blood volume. In the intraoperative bleeding scenario, FFP administration started at 164 min when bleeding volume was 0.42 times total blood volume. In the traumatic bleeding scenario, FFP administration started at 170 min when bleeding volume was 1.63 times the total blood volume. During FFP transfusion, the appropriate PRBC:FFP ratio was 1:0.47 in both scenarios.

PC administration was started at 219 min when bleeding volume was 2.24 times the total blood volume in the intraoperative bleeding scenario and at 225 min when bleeding volume was 2.20 times the total blood volume in the traumatic bleeding scenario. After the start of PC transfusion, the appropriate PRBC:FFP:PC ratio was 1:0.35:0.39 in both scenarios.

DISCUSSION

Computer simulations regarding dilutional coagulopathy have been previously studied [8,9,16]. Here, we designed two separate scenarios and compared them. In the traumatic bleeding scenario, the amount of bleeding per minute reduced by decreased blood pressure compared to intraoperative scenario; as a consequence, the onset of dilutional coagulopathy was delayed. Similar results have been
shown by Hirshberg et al. [8,18].

The key Hirshberg model equations and parameters were not complete as is, and did not work when entered into the STELLA program. Therefore, we modified some parts of the equation to implement the model and attempt to reproduce the results of the Hirshberg study. As such, our results have some differences when compared with Hirshberg, which suggest two options for giving FFP before the prothrombin time (PT) crossover time. PT crossover time means the point during a simulation when a clotting test first crosses its respective threshold. In other words, the time when the plasma fraction goes below 30% of the original plasma fraction. The first option is to use an aggressive lower PRBC/FFP replacement ratio such as 3:2 and the second option is to give two units of FFP concurrently with the first units of PRBCs at the beginning of the operation. This suggestion by Hirshberg et al. [8,18] is a much lower PRBC/FFP ratio than the 5:2 to 5:3 ratio used in many massive transfusion protocols and also much lower than our simulation results of 1:0.47 = 5:2.35 ratio.

Fig. 2. Model prediction of hematocrit (%), plasma dilution (%), bleeding fraction and adjusted platelet count (/ml) in the intraoperative bleeding scenario. Phase 1: bleeding started, but only crystalloid fluid was given to maintain normal blood pressure. Phase 2: PRBC administration was started at 43 min and the bleeding fraction was 0.44. PRBCs were administered. Phase 3: FFP was started at 164 min and the bleeding fraction was 1.67. PRBCs and FFP were given. Phase 4: PC was started at 219 min and the bleeding fraction was 2.24. PRBCs, FFP, and PC were given. Bleeding fraction: cumulative blood loss / initial blood volume. PRBC: packed red blood cells, FFP: fresh frozen plasma, PC: platelet concentrate.

Table 2. The Outcome of Two Scenarios

<table>
<thead>
<tr>
<th></th>
<th>Intraoperative bleeding scenario</th>
<th>Traumatic bleeding scenario</th>
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<tbody>
<tr>
<td>PRBC:FFP</td>
<td>1:0.47</td>
<td>1:0.47</td>
</tr>
<tr>
<td>PRBC:FFP:PC</td>
<td>1:0.35:0.39</td>
<td>1:0.35:0.39</td>
</tr>
<tr>
<td>Time of PRBC administration after initiation of bleeding (min)</td>
<td>43</td>
<td>52</td>
</tr>
<tr>
<td>Time of FFP administration after initiation of bleeding (min)</td>
<td>164</td>
<td>170</td>
</tr>
<tr>
<td>Time of PC administration after initiation of bleeding (min)</td>
<td>219</td>
<td>225</td>
</tr>
<tr>
<td>Bleeding fraction at initiation of PRBC transfusion</td>
<td>0.44</td>
<td>0.42</td>
</tr>
<tr>
<td>Bleeding fraction at initiation of FFP transfusion</td>
<td>1.67</td>
<td>1.63</td>
</tr>
<tr>
<td>Bleeding fraction at initiation of PC transfusion</td>
<td>2.24</td>
<td>2.2</td>
</tr>
</tbody>
</table>

In the Hirschberg model, patient initial bleeding rate was 135 ml/min with 3,297 ml lost and 67% of the estimated blood volume at the beginning of the operation. However, in our traumatic bleeding scenario, initial bleeding rate was 50 ml/min with 1,175 ml lost and 41% of the estimated blood volume, and FFP transfusion was started during the 11th PRBC transfusion. We think the reason why this difference occurred was because the increased massive bleeding without hemodilution incurred a greater loss of clotting factor. Comparing our intraoperative bleeding scenario (FFP transfusion started at 1.67 blood volume loss) with the trauma scenario (FFP transfusion started at 1.63 blood volume loss), dilutional coagulopathy began at less blood loss. Early FFP transfusion in the Hirshberg model could be due to early massive blood loss in that scenario [8].

Hirshberg ignored plasma contained in PRBCs and PC. PRBCs contain a small amount of plasma (30–60 ml) as does PC (approximately 80 ml) [20]. In our model, we assumed PRBCs contain 58 ml of plasma and PC contains 41 ml of plasma, both without CPDA-1 solution, by using information of the blood compartment obtained in the 2013 transfusion guidelines. These are not negligible volumes compared with 124 ml of plasma contained in FFP without CPDA-1 solution. If plasma contained PRBCs and PC is not considered, the calculated PRBC/FFP ratio will decrease. Therefore, the Hirshberg 3:2 PRBC/FFP ratio could be underestimated.

The Pragmatic, Randomized Optimal Platelet and Plasma Ratios trial was designed to address the effectiveness and safety of transfusing PRBC, FFP, PLT in a 1:1:1 ratio compared with a 2:1:1 ratio in patients with trauma who were predicted to receive a massive transfusion. Among patients with severe trauma and major bleeding, early administration of PRBC, FFP, PLT in a 1:1:1 ratio compared with a 2:1:1 ratio did not result in significant differences in mortality at 24 h or 30 days. However, more patients in the 1:1:1 group achieved hemostasis and fewer experienced death due to exsanguination by 24 h. Even though there was an increased use of FFP and PLT transfused in the 1:1:1 ratio.
group, no other safety differences were identified between the two groups [21]. Conversely, other studies have report-
ed beneficial outcomes across a wider range of blood product ratios or goal-directed approaches [22,23].

Our simulation shows a 1:0.47 ratio for PRBC:FFP and a 1:0.35:0.39 ratio for PRBC:FFP:PC, if dilutional coagulopathy is assumed to occur when plasma dilution is < 30%, fibrinogen is < 0.8 g/L, and platelets are < 50,000/ml. Interpreting this result requires circumspection; however, patients generally have a wide range of coagulation problems such as dysfunction of platelets and the coagulation system, increased fibrinolysis, compromise of coagulation by the infusion of colloids, hypocalcemia, disseminated intravascular coagulation-like syndrome, acidosis, hyperthermia, and the destruction of clotting factor in stored FFP [24,25]. In the massive transfusion scenario, avoiding the above listed problems is not easy and computer simulations will not perfectly imitate the conditions of the real body. So, our results can be used only as reference and not indication. Careful observation of operation in the field, communication with surgeons, and repetitive coagulation tests are the most important variables in preventing coagulopathy.

The results of this simulation show that coagulopathy can occur at less blood loss due to the loss of undiluted blood if fluids are not infused after the start of bleeding. In addition, it must be noted that the coagulopathy and transfusion start times were delayed because low blood pressure reduces hemorrhage. Restricting fluid infusion seems to delay dilutional coagulopathy but it can generate circulation problems and can bring about more disastrous consequences to the patient. Therefore, restricting fluid technique to delay reduction of hemorrhage and dilutional coagulopathy must be done cautiously. In addition, the difference in the time when coagulopathy occurred between the two scenarios was negligible in our study. More than likely, the amount of hemorrhage (about 20% of total blood volume) was too small to make a difference and transepipillary refill could have attributed to make the difference even less negligible.

Massive transfusion protocols are widely various among hospitals, with replacement ratios ranging between 10:1 and 5:3 for PRBC:FFP and between 10:6 and 10:12 for PRBC:PLT [26,27]. Clinical suitability and justification of the various massive transfusion practices in trauma or operations is clearly required. Due to the limitations of computer modeling when compared with a biologic system or clinical trial data, our findings surely need validation and further assessment. However, computer simulation models can uncover new predictions and generate data that can be used in place of harmful clinical trials that would be against medical ethics; as such, increased need for computer simulations in clinical experiments is apparent.

In conclusion, according to computer simulation, the appropriate blood component ratio might be a 1:0.47 ratio for PRBC:FFP and a 1:0.35:0.39 ratio for PRBC:FFP:PC, which did not differ between the intraoperative bleeding and traumatic bleeding scenarios.

**SUPPLEMENTARY MATERIALS**

Supplementary data including key model equations and parameters can be found online at https://doi.org/10.17085/apm.20042.

**CONFLICTS OF INTEREST**

No potential conflict of interest relevant to this article was reported.

**AUTHOR CONTRIBUTIONS**


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Management of massive bleeding