

A regional massive hemorrhage protocol developed through a modified Delphi technique

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Abstract

Background: A massive hemorrhage protocol (MHP) enables rapid delivery of blood components in a patient who is exsanguinating pending definitive hemorrhage control, but there is variability in MHP implementation rates, content and compliance owing to challenges presented by infrequent activation, variable team performance and patient acuity. The goal of this project was to identify the key evidence-based principles and quality indicators required to develop a standardized regional MHP.

Methods: A modified Delphi consensus technique was performed in the spring and summer of 2018. Panellists used survey links to independently review and rate (on a 7-point Likert scale) 43 statements and 8 quality indicators drafted by a steering committee composed of transfusion medicine specialists and technologists, and trauma physicians. External stakeholder input from all hospitals in Ontario was sought.

Results: Three rounds were held with 36 experts from diverse clinical backgrounds. Consensus was reached for 42 statements and 8 quality indicators. Additional modifications from external stakeholders were incorporated to form the foundation for the proposed MHP.

Interpretation: This MHP template will provide the basis for the design of an MHP toolkit, including specific recommendations for pediatric and obstetrical patients, and for hospitals with limited availability of blood components or means to achieve definitive hemorrhage control. We believe that harmonization of MHPs in our region will simplify training, increase uptake of evidence-based interventions, enhance communication, improve patient comfort and safety, and, ultimately, improve patient outcomes.

assive bleeding is a leading preventable cause of death following trauma, childbirth and surgery.¹⁻³
There were 5.1 million deaths after traumatic injury worldwide in 2010, mostly affecting young people, accounting for nearly 10% of all deaths.⁴ In the United States, it is estimated that up to 20% of such deaths are the direct result of preventable hemorrhage.⁴⁻⁷ Management of unstable hemorrhagic shock is centred on stabilizing the patient with prompt transfusion of blood components, and rapid identification and treatment of the source of bleeding. Patient outcome is dependent on the availability of rapid definitive surgical intervention, support of a transfusion

medicine and clinical laboratory, prompt access to hemostatic agents and care provided by a high-performing interdisciplinary team.⁸ In the trauma literature, protocolized delivery of massive transfusion streamlines the complexities

Competing interests: For panellists and conflicts of interest, see Appendix 1 (available at www.cmajopen.ca/content/7/3/E546/suppl/DC1).

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of rapid access to surgical care and blood components, decreases variability of treatment, increases protocol compliance, reduces blood component wastage, facilitates interprofessional communication and allows for tracking of metrics for continuous quality improvement.^{9–12}

Most academic institutions have a massive hemorrhage protocol (MHP) in place to rapidly deliver blood components and coordinate care in the setting of traumatic injury.¹³ However, in a recent survey of 150 hospitals in Ontario, the proportion of hospitals with MHPs in nonacademic settings was lower than that in academic settings, and there was significant protocol heterogeneity between hospitals.¹⁴ This was also seen in well-established trauma centres in the US,15 where civilian implementation is still highly variable. 12 Compliance with MHPs is also highly variable, 12 and low compliance may have ramifications for patient outcomes. 16 A systematic review of before-after studies comparing patients with trauma managed in periods with and without an MHP showed an association between MHP use and better survival.¹⁷ The benefits of an MHP have not been tested in rigorous prospective randomized trials. Maintaining high levels of compliance with MHPs appears to be a universal challenge. 12,16 Ontario has the largest population of all the Canadian provinces and provides hospital care in a diverse mixture of settings with high and low health care resources. Massive hemorrhage is an infrequent event in many areas served by small regional hospitals. As a result, access to blood components, laboratory tests of hemostasis and surgical expertise are highly variable. Instituting an adaptable MHP for the province based on the local health care resource setting is needed to streamline the complex logistics of rapid delivery of blood components, facilitate rapid patient transfer where required and reduce the cognitive burden on bedside clinicians. With the ultimate goal of developing such a protocol, we aimed to identify the key principles and quality indicators required to develop a provincial standardized evidence-based MHP template for hospitals.

Methods

Steering committee and panellists

A steering committee composed of transfusion medicine specialists and technologists, and trauma physicians selected the panel members, organized the MHP forum, administered the surveys and analyzed the results. The steering committee assembled a panel of 36 content experts to represent relevant stakeholders. The panel members were selected through the Ontario Regional Blood Coordinating Network, which oversees transfusion use, audits of practice and educational initiatives for the region. The steering committee selected members for their broad expertise and responsibility in administering MHPs across Ontario's geographically diverse network. Panel participation was voluntary and not financially remunerated. Travel reimbursement and meals were provided, and authorship was afforded only to those who participated in all phases of the Delphi process. The panel included anesthesiologists, trauma surgeons, obstetricians, hematologists, transfusion medicine physicians, emergency physicians, prehospital and transport medicine physicians, intensivists, blood supplier representatives, nurses, technologists and a patient. The group represented the geographically diverse health care programs in the province, with representatives from academic hospitals, specialized pediatric institutions, suburban hospitals and smaller, rural hospitals.

The panellists were informed of the purpose and scope of the exercise, and of the requirement to attend a 2-day MHP forum in Toronto and complete all rounds of the Delphi exercise. To standardize the knowledge base of panellists, they were provided with copies of original papers selected by the steering committee and reflecting the most up-to-date evidence in the area of massive hemorrhage management before the first round. The MHP forum had both didactic and interactive sessions, in which content experts reviewed each area of the MHP for 15 minutes, after which there was a 15-minute discussion period for the attendees.

Study design

We chose a modified Delphi technique to establish consensus and the framework for the provincial MHP toolkit. The Delphi technique is a systematic, interactive method that relies on a panel of experts to converge on consensus statements following a series of iterative surveys. 18 Rounds of surveys are continued until consensus is achieved. The Delphi technique is deemed a relevant source of evidence in health care research and is particularly important if randomized controlled trials are unavailable to set health care policies.¹⁹ Our group used this method previously²⁰ and noted its ability to costeffectively include a large number of participants who are geographically dispersed, while preventing unequal representation or biasing of opinions in the consensus through iterations of anonymous voting.21 We modified the Delphi technique to allow for open forum of discussion in round 1 only, to ensure that the broadest range of expert experience was captured in the consensus.

The modified Delphi rounds were conducted independently by each panellist via an emailed survey link (Lime-Survey). Survey responses were anonymized before centralized review. Each statement was independently rated on a 7-point Likert scale from "Definitely should not include" to "Definitely should include" in the MHP, with cut-offs based on previously established Delphi criteria. ²⁰ There was an "opt out" option for each statement to account for possible lack of expertise in a specific area ("Unable to rate as outside area of expertise"). Panellists were asked to provide suggestions to enhance statement clarity with each numerical ranking. Panellists were instructed to answer questions on the basis of what they considered optimal patient care or best practices, rather than what they believed was currently operationally feasible at their institution.

The first round consisted of 43 statements and 8 quality indicators that had been drafted by the steering committee based on available literature and existing MHPs. The



statements were based on the core tenets of existing MHPs, such as rapid and reliable access to blood components, tertiary care transfer and use of antifibrinolytic agents, and the quality indicators were based on quality measures established by the National Surgical Quality Improvement Program. The first round was conducted 1 month before the MHP forum. The survey responses of the first round were anonymized and presented to the panellists and the steering committee on day 2 of the MHP forum for discussion of the phrasing and structure of statements scoring below a median Likert score of 5.5 (see a priori criteria below), as the modification to the Delphi methodology. Round 1 provided panellists with the ability to add statements and quality indicators. To ensure nonbiased and independent statement review, panellists were not provided the ratings and survey comments of the other panel members. After round 1, no further inperson meetings or formal interactions between panellists took place.

Criteria for disposition of items

A priori criteria for disposition of the items in the first round were established as follows:

- 1. Items receiving a median Likert score of at least 5.5 (out of 7) would be accepted as written and not subject to further rounds. These statements were to be incorporated into a provincial MHP as written, unless a clear improvement in phrasing was suggested by a panellist that would not change the intent of the item.
- Items with a median score of 2.6 to 5.4 were to be discussed at the in-person meeting with all the panellists. Following discussion, the items would be revised by the steering committee and sent out electronically in the second round.
- Items with a median score of 2.5 or less were to be removed from further rounds, unless there was strong opposition by the panel, in which case a revision would be drafted for the second round.
- 4. Panellists were provided with a comment box to allow for addition of novel statements and quality indicators on the first round and were allowed to suggest additional statements and quality indicators at the in-person meeting. No additional statements were added after round 2.

A priori criteria for disposition of the items in the second or later round were established as follows:

- 1. Items receiving a median Likert score of at least 5.5 would be accepted as written and not subject to further rounds, unless a clear improvement in phrasing was suggested by a panellist that would not change the intent of the item.
- 2. Items with a median score of 2.5 to 5.4 were to be rewritten on the basis of comments by the panellists and sent out in the third or subsequent round.
- 3. Items with a median score of 2.4 or less were to be removed from further rounds of scoring.
- 4. Where suggested by panellists and/or the steering committee, merging or division of statements could occur where appropriate.

After consensus was reached in the final Delphi round, the statements were circulated via email to the medical directors of transfusion medicine responsible for 150 of the 262 Ontario hospitals with licensed transfusion laboratories. The accompanying letter included a request to distribute the statements to members of their hospital transfusion committees and MHP leaders for feedback. These 304 people represented the clinical and technical leads of the laboratories. All hospitals in Ontario able to issue red blood cells have a medical director who is registered with the provincial Ministry of Health and Long-Term Care, which provided us with an upto-date email contact list. Feedback was collated and recommendations were incorporated by the steering committee where necessary to improve the clarity of the statements and their justification.

The initial drafting of the consensus statements was completed by March 2018. Round 1 of the consensus panel was completed on Apr. 13, 2018, and was discussed in person on Apr. 21, 2018 in the MHP forum. Rounds 2 and 3 were subsequently completed in June 2018 and August 2018, respectively.

Ethics approval

Ethics approval was not required for this Delphi exercise.

Results

Of the 44 experts invited to participate, 36 agreed (Appendix 1, available at www.cmajopen.ca/content/7/3/E546/suppl/DC1). A 100% response rate was achieved from panellists in all 3 rounds of the modified Delphi exercise. The progression toward consensus based on Likert scores is shown in Figure 1, and the overall degree of consensus for each statement by the last round is shown in Figure 2. Consensus was achieved for all statements by round 3, yielding a final consensus document with 42 statements and 8 quality indicators. Owing to lack of consensus on round 1, statements 7, 22, 23 and 34 were broken into their individual components for rounds 2 and 3 to assist with understanding the components driving the lack of consensus (Figure 1). Unless otherwise specified, all statements and quality metrics also apply to a pediatric MHP. The statements together with a brief rationale for each are presented in Table 1, in a logical order rather than in order of clinical importance.

Four areas required additional rounds and major modifications: 1) selection of the name of the protocol, 2) selection of the laboratory resuscitation targets, 3) determination of the pack configurations and 4) clarification of the role of recombinant factor VIIa. The primary obstacle to selecting a name for the protocol was that many hospitals already had long-standing MHPs with specific names. Consensus on the laboratory targets and pack configuration was achieved in the third round by splitting statements into subsections. The recombinant factor VIIa statement required 3 rounds of review to ensure that the phrasing mitigated the panellists' apprehensions regarding this controversial therapy. Following the generation of consensus statements, no



Statement	Round 1: median score (range; abstaining votes)	Round 2: median score (range; abstaining votes) or counts where appropriate	Round 3: median score (range; abstaining votes) or counts where appropriate
2	7 (5–7; 1) 7 (5–7; 1)		
3	7 (4–7; 3)		
4	7 (1–7; 1)		
5	6 (2–7; 2)	7 (4–7; 3)	
6	7 (5–7; 1)	. (, 0)	
7	6 (2–7; 2)	6 (1–7; 1)	Overhead "Code Transfusion": rank 1
8	6 (3–7; 1)	2 (* 1, 1)	
9	7 (4–7; 2)		
10	6 (2–7; 4)	7 (5 7 4)	
	7 (4–7; 4)	7 (5–7; 1)	
11	7 (4–7; 1)		
12	7 (3–7; 1)		
13	7 (3–7; 1)		
14	6 (2–7; 4)	7 (2–7; 1)	
15	7 (3–7; 2)		
16	7 (3–7; 1)		
17	7 (1–7; 1)	7 (2–7; 1)	
18	7 (4–7; 2)		
19	NA	7 (1–7; 0)	
20	7 (4–7; 2)		
21	6 (2–7; 1)		
22	7 (2–7; 1)	"Yes" to need for test: • CBC, INR, iCalcium 36/36; fibrinogen 35/36; pH 34/36 • aPTT 29/36; electrolytes 32/36; lactate 30/36	
23	7 (2–7; 3)	 Hemoglobin > 80 g/L 31/36 INR < 1.8 26/36 Fibrinogen > 1.5 g/L 24/36 Platelets > 50 x 10⁹/L 26/36 iCalcium > 1.15 mmol/L 17/36 INR < 1.5 11/36 Fibrinogen > 2.0 g/L 12/36 Platelets > 100 x 10⁹/L 12/36 iCalcium > 1 mmol/L 11/36 	
24	7 (4–7; 5)	7 (5–7; 4)	
25	7 (4–7; 4)	. (0 ., .)	
26	7 (4–7; 7)		
27	7 (2–7; 4)		
28	6 (4–7; 3)	7 (2–7; 1)	
29	7 (2–7; 5)	. (= ., ./	
30	7 (4–7; 1)		
31	7 (2–7; 1)	7 (6–7; 1)	
32	7 (2–7; 11)	7 (5–7; 5)	
33	7 (1–7; 3)	() , , , ,	
34	6 (3–7; 4)	6 (1–7; 4)	Platelet transfusion based on count: 7 (1–7; 4) Communicate if no platelet transfusion: 6 (2–7; Box 1: 7 (1–7; 2) Box 2: 7 (2–7; 4) Box 3: 7 (1–7; 4) Small hospital box 2: 7 (2–7; 5) Convert to laboratory-guided: 7 (2–7; 5)
35	7 (3–7; 8)	7 (2–7; 6)	
36	7 (4–7; 6)	7 (5–7; 6)	
37	6 (1–7; 11)	7 (3–7; 4)	
38	7 (1–7; 1)	7/4 7 4	
39	7 (1–7; 2)	7 (4–7; 1)	
40	6 (4–7; 5) 6 (2–7; 8)	7 (4–7; 4)	
	6.5 (2–7; 5)		
11	7 (5–7; 1)		
12-1	7 (1–7; 4)	7 (4–7; 2)	
12-2	7 (1–7; 2)	7 (2–7; 1)	
12-3	6 (1–7; 5)	7 (5–7; 1)	
	7 (1–7; 6)	7 (4–7; 1)	
	6 (1 7, 0)	7 (1–7; 1)	
42–5	6 (1–7; 2)		
42-5 42-6	6.5 (1–7; 2)	7 (4–7; 1)	
42-4 42-5 42-6 43-7 42-8			

Figure 1: Median scores on a 7-point Likert scale for the 3 rounds of the Delphi exercise. Counts are provided for cases in which panellists were asked to rank or choose between options rather than use the Likert scale. Green = passed with minor or no phrasing adjustment (empty green box = passed on a preceding round); red = statements that did not pass based on numerical scoring and/or critical written comments, resulting in a major content revision; gray = statements that were not accepted and were then merged for the subsequent round; blue = a new statement that was added after round 1. Split rows denote the need for division of the statement into its components for the scoring round. Note: CBC = complete blood count, iCalcium = ionized calcium, INR = international normalized ratio, NA = not applicable, aPTT = activated partial thromboplastin time.



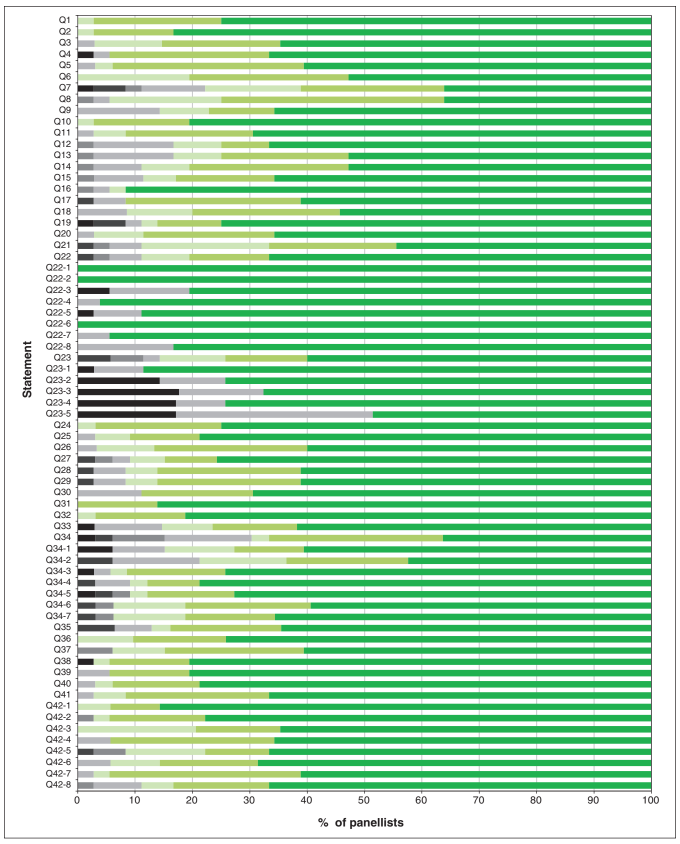


Figure 2: Proportion of Likert scores at the time of the final consensus round for each statement. Panellists were asked to indicate agreement on a 7-point Likert scale (7 = highest level of agreement). Black and gray = scores less than 5; green shades = scores of 5 or more. Responses of "no," "uncertain" and "yes" are denoted as black, gray and green, respectively.



substantive changes to the statements were made, but modifications were required to clarify the supporting text.

The external review by the medical directors of transfusion did not necessitate changes to the statements; however, the

feedback provided valuable input to strengthen the supporting text and provide input on logistical and implementation challenges. All statements and the contents of the manuscript were approved by all authors and panellists.

Statement	Rationale
All hospitals shall have a protocol to guide the management of a massively bleeding patient.	The panel concluded that an MHP is required to standardize the approach to the massively bleeding patient for all hospitals. For the purposes of the MHP, a hospital is defined as any organization that either maintains a red blood cell inventory or staffs an emergency department, urgent care centre, critical care unit, labour and delivery unit, or operating room. The panel recognized there are small clinic facilities where a bleeding patient may be encountered but where transfusion is currently not available and an MHP would not be appropriate. The panel concluded that a policy for rapid transport of patients with massive hemorrhage to a facility with an MHP would be required at such a facility.
2. The protocol shall be developed by a multidisciplinary team and approved by the hospital transfusion committee (or other relevant multidisciplinary committee).	The MHP requires support from multiple hospital services including, but not limited to, emergency, trauma, surgery, anesthesiology, critical care, blood transport personnel, communication services and laboratory personnel. The protocol should be reviewed and approved by the hospital transfusion committee (or other relevant hospital committee) and the medical advisory committee.
3. The protocol shall incorporate the principles of damage-control resuscitation, specifically giving highest priority to treating the source of hemorrhage.	Damage-control resuscitation principles in traumatic injury include abbreviated surgical and/or endovascular interventions for hemorrhage control and management of intra-abdominal contamination, and critical care support to correct deranged physiologic measures (hypothermia, acidosis, coagulopathy), with definitive surgical repair delayed until stabilization and hemostatic control have been achieved. ²² In the severely injured trauma population, damage-control resuscitation is associated with reduced mortality, although the approach has never been tested in a randomized controlled trial. ^{1723,24} Ongoing hemorrhage leads to worsening coagulopathy and other physiologic derangements. ²⁵ Although the role of damage-control resuscitation outside of traumatic injury is unknown, prompt hemorrhage control is likely to be an important component of care. ^{26,27}
4. The protocol shall consider the available resources at the institution.	The hospital must consider the available resources of the institution when developing the local protocol. Centres caring for pediatric patients should ensure personnel are prepared for weight-based dosing and the use of size-specific equipment (e.g., warming devices, intravenous infusion equipment). Smaller and more remote hospitals located at a distance from the blood supplier will need to make adjustments to streamline their MHP to compensate for the limited number of team members, blood component inventory and laboratory testing menus, and the ability to provide definitive surgical or endovascular control of hemorrhage. The MHP will need to specify, if required, which patients should be transferred in a timely manner to other facilities for definitive treatment and how this should be achieved. Examples for simplification for smaller/remote sites include 1) prelabelled uncrossmatched RBC units ready for immediate transfusion, 2) preprepared laboratory sample collection kits, 3) administration of a single bolus of tranexamic acid rather than an infusion, 4) administration of PCCs and fibrinogen concentrate instead of plasma and cryoprecipitate, 5) use of point-of-care technology for laboratory testing and 6) cross-training hospital personnel from other patient care areas.
5. A single protocol for all patients is preferred in order to ensure compliance; there should be specific guidance provided for selected patient populations (e.g., obstetrical patients should receive early fibrinogen replacement).	A survey from academic hospitals showed that 60% of respondents have a single protocol for all patients. Compliance with a single MHP is poor in published studies, 12,16,29 which raises the concern that consistent care would be further compromised by multiple protocols for different bleeding scenarios. The panel recommended a single, standardized protocol in response to the massively bleeding patient with options to tailor the protocol for specific patient populations. For example, in massive obstetrical hemorrhage, consideration should be given to measuring fibrinogen levels early and repeatedly, administering fibrinogen replacement if the level falls below 2.0 g/L ³⁰ and using an intrauterine balloon device as a bridge to definitive bleeding control. In gastrointestinal hemorrhage, consideration should be given to prompt endoscopic therapy for hemorrhage control. And hemorrhage after cardiac surgery, there is evidence to support the use of viscoelastic testing (as compared to standard laboratory tests) in reducing the risk of major bleeding. Pediatric patients require weight-based dosing of blood components and hemostatic adjuncts, consideration of potentially higher transfusion triggers depending on comorbidities and age, and provider awareness of increased risk or hyperkalemia and hypothermia.
6. The protocol should be reviewed at a minimum of every 3 years.	The science and clinical trial activity in the area of massive hemorrhage, coagulopathy and MHPs are rapidly evolving. Each institutional MHP should be reviewed at a minimum of every 3 years to ensure alignment with the scientific evidence and the provincial MHP. The protocol revision should be conducted by a multidisciplinary team as detailed in statement 2, and approved by the hospital transfusion committee and medical advisory committee.



Table 1 (part 2 of 7): Massive hemorrhage protocol statements and rationale

Statement Rationale

7. The protocol shall be called "The Massive Hemorrhage Protocol" and, if activated as an overhead announcement, referred to as "Code Transfusion."

The existence of several different terms for the protocol across Ontario has created confusion and delays to activation (e.g., a trainee calling communications to activate the Code Omega protocol at a hospital that activates the protocol by calling the transfusion medicine laboratory to activate the "massive transfusion protocol"). The panel, after much deliberation, chose the protocol name "Massive Hemorrhage Protocol" for the following reasons: 1) massive transfusion is most commonly defined in adults as a transfusion of 10 or more units of RBCs in a 24-hour period; however, some patients will not survive to receive 10 units, and many patients who receive 4-10 units need additional therapies contained in an MHP; 2) the name highlights the importance of definitive hemorrhage control; and 3) an MHP is more than just a transfusion protocol and includes nontransfusion interventions (e.g., maintenance of normothermia, use of antifibrinolytics). The panel agreed that the method for MHP activation should be site-specific and be clearly defined in the protocol but that if a hospital-wide overhead announcement was implemented, a standard term should be used at all institutions. The consensus term chosen by the panel is "Code Transfusion" owing to its clarity, ease of pronunciation and lack of overlap phonetically with other "colour" codes (e.g., Code Bleed or Code Blood with Code Blue). The value of an overhead announcement is that it provides redundancy if the paging system fails and notifies all hospital employees that the laboratory is under acute pressure (and to refrain from calling for nonemergency blood products and nonurgent test results).

8. Participating team members should have access to formal training and drills to increase awareness, adherence and effective delivery of the MHP.

To achieve high levels of team performance and protocol adherence, team members require access to formal training material and exposure to multidisciplinary drills or simulations. This is particularly important for high-stress and rarely encountered massive hemorrhage scenarios. Simulations have been successfully employed for training in obstetrical hemorrhage,³⁹ pediatric hemorrhage⁴⁰ and trauma.⁴¹ A systematic review of 33 studies involving 1203 resident and medical student participants showed that simulation was associated with improved provider behaviour and patient outcomes.⁴² In a systematic review of 13 studies of trauma team training, both nontechnical skills and team-based performance improved.⁴³ Importantly, these improvements extend to patient outcomes, as simulation-based training is associated with improved outcomes in trauma and cardiac arrest care.^{44,45}

9. The written MHP should be readily accessible as a reference tool for all team members.

To achieve high levels of protocol compliance among staff, ready access to the MHP is required. The local institution should develop resources (either in electronic or paper format) to assist clinicians with MHP compliance. The format and medium should be dictated by the local hospital circumstances.

10. The transport service(s) should be promptly notified if the decision is made to transfer the patient to another hospital for definitive hemorrhage control. If required, the patient should be transferred as soon as and as safely as possible by appropriate staff and transport resources, to an institution where definitive hemorrhage control can be performed.

There are 150 hospitals in Ontario that have access to transfusion support. Owing to Ontario's large geographic size and numerous remote regions, it would not be possible to have large stocks of blood components available at all hospitals without very high levels of wastage. Timely evacuation of massively bleeding patients from smaller centres to larger centres capable of definitive hemorrhage control is needed for 2 reasons: 1) the small blood stocks held in remote hospitals (typically a small number of RBCs, no platelet pools and limited stocks of frozen plasma) and 2) lack of access to definitive surgical or radiologic intervention to allow for hemorrhage control. There is little published on evacuation time targets within civilian settings. Rapid evacuation (< 60 min) among military patients with trauma who had noncompressible torso injury and amputation injury was associated with reduced mortality. 46 Clinicians working with limited capacity to achieve surgical hemostasis should aim to transfer as soon and as safely as possible.

11. The protocol shall have activation criteria.

Underactivation (i.e., delayed or no activation of MHP for patients who require hemorrhage control and blood components) could be catastrophic, as it may result in otherwise preventable exsanguination. A retrospective study suggested that delay in initial blood component administration is associated with worse outcomes (each 1-min delay to the arrival of the first pack of blood components is associated with a 5% increase in the risk of death). 47 In contrast, overactivation (i.e., MHP activation that is ultimately not required) may lead to unnecessary transfusion, wastage of blood components and diversion of human resources away from competing needs. Despite concern that appropriate and timely activation are critical, there are no criteria with both high sensitivity and high specificity for predicting the need for massive transfusion. The 2 most commonly used scores validated in this setting are the Shock Index (blood pressure divided by heart rate or modified pediatric Shock Index⁴⁸) and the ABC score (1 point each for penetrating injury, blood pressure ≤ 90 mm Hg, heart rate ≥ 120 beats/min and positive results of FAST [Focused Assessment with Sonography for Trauma]), with the Shock Index performing slightly better than the ABC score in traumatic injury.⁴⁹ New data suggest that resuscitation intensity (≥ 4 units of fluid in the first 30 min, with 1 unit defined as any of 1 unit of RBCs, 1 unit of plasma, 500 mL of colloid or 1 L of crystalloid) may represent an important alternative metric to identify patients who require MHP activation.⁵⁰ In pediatric patients, a retrospective study of combat-injured children defined massive transfusion as the requirement for 40 mL/kg or more of blood components transfused within 24 hours.⁵¹ Given the current lack of evidence to support one set of activation criteria over another, the activation criteria should be set by the hospital to meet the needs of the local patient population.



Table 1 (part 3 of 7): Massive hemorrhage protocol statements and rationale					
Statement	Rationale				
12. The protocol shall have termination criteria.	Termination of the protocol allows personnel to return unused blood components to regular inventory, cease ordering blood components from the blood supplier, cease thawing of frozen components and divert resources to competing needs. In contrast, premature termination may lead to a reduction in the number of team members at the bedside, in the frequency of laboratory testing and in the availability of blood components. Termination should be considered when bleeding source control has been attained, hemodynamic stability has been achieved, vasopressor requirements have diminished, and the transfusion rate has slowed such that additional transport personnel are no longer required. Typically, when these features are present, transfusion decisions can be guided by laboratory test results. ⁵² As no explicit criteria have been validated, termination criteria should be determined at the local hospital level. The method to communicate the termination of the MHP should be specified in the local hospital protocol.				
13. The protocol shall specify the team members required to respond when the protocol is activated.	Executing all the necessary tasks specified in an MHP, in addition to all the other clinical tasks required to achieve surgical control of blood loss, will require mobilization of an interdisciplinary team. The precise composition of the clinical team can be modified according to the acuity of the hemorrhage, the location of the patient, the type of hemorrhage and the institution's available resources. For example, the neonatal team will be required to attend postpartum hemorrhages to provide immediate care for the neonate, whereas in trauma MHPs managed in the trauma room, where nursing to patient ratios are already high, additional nursing staff may not be required. Given the association between survival and the time of arrival of the first cooler of blood components, a dedicated transport team for both blood samples and components is critical.				
14. The protocol should specify how the lead clinician at the bedside is designated.	How the lead clinician for the MHP is assigned should be specified in the local hospital protocol as it will be highly variable depending on the patient population served and the institutional resources. A broad range of physicians could serve as the team leader. In addition, in smaller organizations without on-site physicians, a nurse practitioner or midwife may be the most appropriate team leader. There may be a transition in leadership as the patient moves from one location to another. The process of handover from one leader to the next should be explicitly stated in the protocol. There must be training in nontechnical skills for the team leads to promote high performance for communication, situational awareness and decision-making skills. In simulation training, higher performance on nontechnical skills by the team lead (situational awareness and decision-making) correlates with critical task completion and improved team performance. ⁵³ Simulation training for clinicians leading trauma resuscitation improves confidence and reduces anxiety. ⁵⁴ The provision of formal feedback to trauma team leaders in training by faculty is associated with improvement in leadership skills over time. ⁵⁵				
15. The protocol shall specify the team member(s) designated to be responsible for blood component and sample transport.	The protocol shall specify the team members designated to be responsible for both the transportation of blood components and patient blood samples for laboratory testing. Although the protocol specifies the use of ratio-based resuscitation (standardized RBCs to plasma) to mitigate the risk of coagulopathy, this does not prevent overtransfusion or provide assurance that coagulation competence will be maintained. Early and repeated laboratory testing (with rapid transportation of the samples to the laboratory) to confirm adequacy of transfusion resuscitation is required. It is also critical that blood components be rapidly supplied to the bedside and that empty coolers be returned to the transfusion medicine laboratory.				
16. The transfusion medicine laboratory and the core laboratory shall be notified of all MHP activations.	Early and prompt notification of the transfusion medicine laboratory will assist with timely blood component delivery, rapid transition to group-specific blood and designation of the transfusion medicine technologist team leader. A single individual on the clinical side should be the sole source of contact between the clinical team and the transfusion medicine technologist leader so as to reduce the risk of duplicate transfusion orders. Activation of the core laboratory technologists will ensure designation of the laboratory technologist team leader, rapid identification of MHP samples, prioritization of the testing, complete testing of all required tests for the MHP and immediate communication of test results to the clinical team.				
17. All critical laboratory results and important coagulation parameters (hemoglobin, platelet count, INR and fibrinogen) shall be communicated verbally to the clinical team as soon as they are available.	During MHP activation, the clinical team may not have ready access to the electronic health record owing to patient acuity and clinical area layout. It is therefore required that all critical results (preliminary or complete, and as defined by the local laboratory) and important coagulation results (hemoglobin, platelet count, INR and fibrinogen) be verbally communicated to the clinical team as soon as the results are available. This may mitigate the risks of undertransfusion or overtransfusion, and improve time to correction of other biochemical derangements (hyperkalemia, hypocalcemia, acidosis). The "push of information" is thought to be an important tool to improve team performance. Consideration should be given to having dedicated mobile phones to mitigate the risk of communication failure between the laboratory and the clinical team due to rapid movement of the clinical team from one hospital location to another.				
18. The timing of protocol activation and termination shall be recorded in the patient's chart.	Documentation of the activation and termination times must be recorded in the patient chart in the format specified by the local institutional policy. This could be documented by hand or electronically in the nursing or physician notes, or in the electronic computerized physician ordering system. These times are necessary during the review of the patient chart for the purposes of quality improvement.				



Table 1 (part 4 of 7): Massive hemorrhage protocol statements and rationale

Statement Rationale

19. Patients and/or their substitute decision-maker for whom the massive hemorrhage protocol was activated should be informed.
Actual (e.g., transfusion-associated circulatory overload, hyperkalemia) and potential adverse effects should be disclosed. Furthermore, women of child-bearing potential should be informed of the risk of red blood cell alloimmunization.

At the earliest possible opportunity, the most responsible physician (or delegate) must have a conversation with the patient and/or the substitute decision-maker regarding why the MHP was activated, the number and types of components transfused, the transfusion complications observed and the potential long-term consequences of transfusion. Informed consent for transfusion should be obtained as per local hospital policy. Patients have variable perceptions related to transfusion risks,⁵⁸ and accurate communication of the potential risks is important to achieve patient-centred care. Women of child-bearing potential should be informed of the risk of red blood cell alloimmunization, which may result in hemolytic disease of the fetus and newborn, and should be counselled to undergo red blood cell antibody screening 6 weeks and/or 6 months after transfusion (many antibodies are evanescent, and there is a brief window for detection).⁵⁹

20. The collection and testing of the group and screen sample shall be prioritized in the protocol to mitigate the impact on group O red blood cells and AB plasma stocks. Both group O RBCs and AB plasma are in chronic short supply in Canada. The proportion of group O RBCs transfused to non–group-O recipients is increasing, with trauma accounting for 10% of this pressure on group O blood stocks. ⁶⁰ The vast majority of AB plasma units are transfused to non-AB recipients. ⁶¹ Given the pressure on AB plasma stocks, it has not been possible to provide male-only AB plasma for all recipients, with resultant cases of transfusion-related acute lung injury from female AB plasma. ⁶² Hence, the draw of the group and screen sample, rapid transport of the sample to the laboratory and testing of the sample should be prioritized.

21. Laboratory testing should be done at baseline and at a minimum hourly until the protocol is terminated.

See rationale for statement 22.

22. The recommended minimum laboratory testing (where the test is available) at each blood draw should be: CBC, INR, activated partial thromboplastin time (aPTT; baseline only), fibrinogen, electrolytes, calcium (ionized), blood gas (pH and base excess) and lactate

Baseline laboratory testing is prognostic, 63 identifies patients receiving oral anticoagulant therapy in need of reversal and directs immediate need for components in excess of the base ratio of RBCs to plasma. Although the proposed MHP includes the use of early ratio-based resuscitation for plasma before laboratory test results of coagulation are available, this does not guarantee that coagulopathy will be prevented and raises the risk of transfusion of unnecessary blood components. Laboratory confirmation of adequate hemostatic resuscitation is required at least hourly. Current guidelines recommend early and repeated measures of hematology and coagulation parameters.²⁴ The measurement of aPTT is recommended only at baseline to detect the anticoagulant effect of certain anticoagulants (e.g., dabigatran) and preexisting bleeding disorders (e.g., hemophilia). If the baseline INR and aPTT are correlated, further aPTT measures are not indicated and may in fact delay release of the other coagulation test results.⁶⁴ Increased aPTT in postpartum hemorrhage is associated with worse outcomes; however, there is considerable overlap and minimal difference between outcome groups, so elevated aPTT is not clinically useful.30 Transitioning from blind ratio-based component therapy to one based on conventional laboratory testing or point-of-care viscoelastic testing has the potential to minimize unnecessary transfusions and allow for targeted component therapy.³⁴ Biochemical tests (e.g., potassium, calcium and pH) may indicate potential complications from massive transfusion or inadequate resuscitation of the patient in hemorrhagic shock. Lactate measurements are also predictive of mortality, although the role of serial measurements in improving patient outcomes has not been confirmed in clinical trials.65

23. The protocol should state the minimum laboratory protocol resuscitation targets for transfusion: 1) hemoglobin > 80 g/L (RBC); 2) INR < 1.8 (plasma or prothrombin complex concentrates); 3) fibrinogen > 1.5 g/L (cryoprecipitate or fibrinogen concentrates); 4) platelets > 50 × 10°/L; 5) ionized calcium > 1.15 mmol/L. Relevant transfusion targets can also be used if viscoelastic testing is performed.

As there are no prospective studies evaluating laboratory resuscitation targets in the setting of massive bleeding, the suggested laboratory targets are based on the consensus opinion of the panellists and are concordant with the published literature. ^{24,66} These are minimum targets to be maintained throughout the resuscitation and are not meant to be overly prescriptive (i.e., restricting blood component issue based on the above values). Certain pediatric populations, such as neonates, those with congenital heart disease, those receiving extracorporeal life support and those in severe respiratory distress, may require higher thresholds for RBC transfusion during an MHP.³⁶⁻³⁸

24. All massively bleeding patients should have a temperature measured within 15 minutes of arrival or protocol activation, and then at a minimum of every 30 minutes (or continuously where available) until the protocol is terminated.

See rationale for statement 26.



Statement	Rationale
25. All patients should receive interventions to prevent hypothermia and achieve normothermia (≥ 36°C).	See rationale for statement 26.
26. All patients should receive warmed intravenous fluids, red blood cells and plasma to avoid hypothermia.	In both traumatic injury and postpartum hemorrhage, temperature monitoring is infrequently performed and, when the temperature is measured, hypothermia is common. ^{67,68} Hypothermia in traumatic injury is associated with worse outcomes, ^{69,70} although prospective trials have not confirmed whether aggressive warming protocols would alter outcomes. ⁷¹ Mild hypothermia is associated with a 22% increase in the risk of transfusion. ⁷² Warming of patients improves their comfort, and, therefore, even in the absence of a confirmed survival benefit, it should be a core part of every MHP. ⁷³
27. Red blood cells should be delivered in a validated container to prevent wastage.	Since RBCs are a valuable resource, strategies to reduce wastage during transport to and storage at the patient bedside are required. Numerous investigators have validated that wastage can be mitigated with appropriate temperature-controlled devices, with resultant substantial cost savings. ^{74,75} At large academic centres with frequent MHP activation, all components should be transported in validated containers to mitigate component wastage.
28. The MHP protocol should ensure there are processes in place to ensure an uninterrupted supply of blood components to the bedside.	The local MHP should include processes to ensure an uninterrupted supply of blood components to the bedside until termination. Specifically, the next cooler should be brought to the patient location before the previous cooler is empty. This will minimize the risk of lacking necessary blood components during the resuscitation. The person assigned to maintain the uninterrupted supply of blood components should be specified in the protocol. The procedure for requesting the next set of blood components should be stated in the protocol, easy to perform in the setting of massive hemorrhage and designed with the intention of preventing wrong-patient transfusion errors. The delivery of blood components to the bedside should not be equated with an order for transfusion.
29. If the blood group is unknown, O Rh D-negative red blood cells should be used only for female patients of child-bearing potential (age < 45 yr).	O Rh D-negative stocks are insufficient to allow all patients of unknown blood group to be supported with O Rh D-negative RBCs until the blood group result is released in the laboratory information system. The risk of alloimmunization in an Rh D-negative patient after exposure to Rh D-positive RBCs in the setting of major bleeding is 20%. ^{76,77} Immunization to the D-antigen is relevant only for females who wish to have future pregnancies. Over 99% of births occur among women less than age 45 years, ⁷⁸ and hospital MHPs should restrict the use of O Rh D-negative RBCs for women less than this age. For conscious women, efforts should be made to determine their age early in the course of care so that the transfusion medicine service can be instructed to supply the optimal Rh D-type of blood. The risk of immunization from Rh D-positive platelets is 1%, and therefore Rh-immunoglobulin should be provided only to Rh D-negative women less than age 45 (after transfer to the intensive care unit but within 72 h of the Rh D-incompatible platelet transfusion). ⁷⁹
30. Uncrossmatched red blood cells shall be available at the bedside within 10 minutes of MHP activation.	See rationale for statement 31.
31. In bleeding patients in need of red blood cell transfusion, uncrossmatched red blood cells should be transfused until crossmatch-compatible red blood cells are available.	In retrospective analyses in trauma resuscitation, faster time to delivery of the first pack of RBCs was associated with superior survival (every 1-min delay to the first pack was associated with a 5% increase in the odds of mortality). Collection of the group and screen sample, transport of the sample to the laboratory, centrifugation of the sample, testing and result release into the laboratory information system require about 70–90 minutes. Therefore, following MHP activation, it is not appropriate to wait for crossmatch-compatible RBCs. The transfusion laboratory must have a protocol and process for the immediate release of uncrossmatched RBCs. In severe traumatic injury, in cases in which communication from the prehospital emergency services suggests the patient will need immediate transfusion owing to hemodynamic instability and severe injury, it is appropriate to order RBCs to the emergency department in advance of patient arrival.
32. There is no threshold of units of group O red blood cells above which a switch to group-specific red blood cells is prohibited. The switch to group-specific red blood cells and plasma should be done as soon as possible.	Each unit of RBCs in Canada is produced with a minimal amount of residual plasma (< 30 mL per unit) and, therefore, even after transfusion of 10–20 units of group O RBCs, the amount of incompatible plasma is trivial and does not preclude a transition to group-specific RBCs.
33. The protocol shall state the reversal strategy for commonly used orally administered anticoagulants.	The MHP protocol shall include a table with all approved anticoagulant therapies and their appropriate reversal strategy, including the dosage(s) of the therapies to be administered.



Table 1 (part 6 of 7): Massive hemorrhage protocol statements and rationale

Statement Rationale

34. The initial management of the rapidly bleeding patient that precludes the use of laboratory-guided transfusion should begin with immediate red blood cell (RBC) transfusion and then transfusions at an RBC:plasma ratio of 2:1.

Two prospective randomized trials failed to confirm a survival benefit of an RBC:plasma ratio of 1:1 (compared to 2:1). 80.81 A large retrospective review of experience before and after implementation of 1:1 resuscitation failed to show a mortality benefit. 82 The Canadian consensus conference on massive transfusion recommended an RBC:plasma ratio of 2:1 followed by transition to laboratory-guided administration of blood components as soon as possible. 83 The standard approach outlined below, which is based on expert consensus, is applicable to most large adult hospitals. No blood components should be transfused without a clear order and specified infusion rate from the team leader or delegate. Simplified options are provided for institutions that do not stock plasma, platelets and/or cryoprecipitate (or are unable to provide the components in a rapid manner owing to limited numbers of personnel or lack of thawing devices), or that cannot provide definitive surgical or radiological care, and the goal is to stabilize in preparation for transport by land or air ambulance. Pediatric institutions should develop age- and weight-based MHP component protocols to ensure that blood components and fractionated coagulation factors are delivered in appropriate ratios. In institutions that care for pediatric patients, the transfusion boxes must come with clear instructions for the clinical team to mitigate the risk of over- or undertransfusion.

Standard approach

- 1. Box 1 should contain 4 RBC.
- 2. Box 2 should contain 4 RBC, 4 plasma.
- 3. Box 3* should contain 4 RBC, 2 plasma and fibrinogen replacement (10 units of cryoprecipitate or 4 g of fibrinogen concentrate).
- Platelets, when stocked in the hospital transfusion laboratory, should be transfused based on the platelet count.

Simplified options for smaller organizations

- 1. As per standard approach.
- 2. Box 2 (where plasma **not** stocked in hospital transfusion laboratory) should contain 4 RBC, 2000 IU of PCC and 4 g of fibrinogen concentrate. Efforts should be made to transfer the bleeding patient to a centre capable of definitive hemorrhage control.
- 3. As per standard approach.
- 4. Platelets, when **not** stocked in the hospital transfusion laboratory, should be ordered in for transfusion (if patient cannot be promptly transferred out). If the patient is transferred before platelets are transfused, this should be communicated to the receiving hospital.

35. Recombinant factor VIIa (rVIIa) should be considered only when massive hemorrhage is refractory to surgical hemostasis, medical optimization of coagulation parameters, acidosis and hypocalcemia, and be used in consultation with an expert in the management of coagulopathy in the massively bleeding patient.

Recombinant activated factor VIIa (rVIIa) has not been shown to reduce mortality in prospective randomized controlled trials.^{84,85} In contrast, rVIIa is associated with an increase in thromboembolic complications.⁸⁴ Given the concerns regarding lack of efficacy and potential risks, all other lower-risk hemostatic therapies should be exhausted, and rVIIa should be used only in consultation with an expert in the management of coagulopathy of the massively bleeding patient.

36. Fibrinogen concentrate, 4 g (equivalent to approximately 10 units of cryoprecipitate), can be used as a reasonable alternative to cryoprecipitate for fibrinogen replacement.

Cryoprecipitate in Canada is provided as individual units that must be thawed, reconstituted with saline and then pooled. This takes about 30–45 minutes of technologists' time and may compete with their ability to perform laboratory testing or prepare other components. The product can be kept for only 1 year after donation. It must be transported frozen at all times. Once thawed and pooled, it expires after 4 hours. Given the time-intensive preparation requirements and limited shelf life, it is reasonable for some hospitals to transition to pathogen-reduced fibrinogen concentrates. There are no large randomized controlled trials of cryoprecipitate and fibrinogen concentrates to determine equivalence, although a large trial in cardiac-surgery—related hemorrhage (FIBrinogen REplenishment in Surgery [FIBRES] trial) is ongoing.⁸⁶ For pediatric patients, a dosage of about 50 mg/kg of fibrinogen concentrate up to a maximum of 4 g is suggested.⁸⁷

37. At institutions lacking sufficient resources to issue plasma (e.g., no thawing device or no plasma stocked in inventory), prothrombin complex concentrates (PCCs), 2000 IU, can be substituted for coagulation factor replacement. Fibrinogen replacement should be given concurrently with PCCs unless the fibrinogen level is known to be \geq 1.5 g/L.

Similar to the challenges with cryoprecipitate, some smaller organizations may have challenges in providing plasma during an MHP (no thawing device or not stocked in the laboratory owing to rarity of use). In these situations, a reasonable option is to transfuse PCCs and fibrinogen concentrates. This is a common strategy used in many European countries, and outcomes appear to be similar to those with a plasma resuscitation strategy in trauma, usually guided by viscoelastic point-of-care testing.⁸⁸ This strategy should be seen as a bridge before transport to an institution capable of definitive surgical management and more complete transfusion support. For pediatric patients, a dosage of 25 IU/kg of PCCs (rounded to the closest 500 IU) up to a maximum of 2000 IU is suggested.^{89,90}



Table 1 (part 7 of 7): Massive hemorrhage protocol statements and rationale					
Statement	Rationale				
38. Pretransfusion bedside patient and product identification check shall be performed before transfusion of any component to avoid mistransfusion.	Transfusion-related errors remain common in the emergency department. 91,92 Under no circumstances can the pretransfusion bedside patient and product identification check be aborted, especially in mass-casualty scenarios, where there may be multiple patients receiving blood components simultaneously.				
39. Patient demographics shall not be updated/changed until after termination of the protocol. Once the MHP is terminated, patient demographics should be updated as soon as possible.	Patients admitted during major hemorrhage or after traumatic injury are frequently registered with a temporary name and number (e.g., "Unidentified," "Andrew") or with an incomplete registration (e.g., no date of birth). Modifications to key identifiers during active resuscitation may delay the issue of blood components from the transfusion service or result in an erroneous incompatibility detected at the pretransfusion bedside check. The update of the patient identification should be delayed until the patient's condition has stabilized and with coordination between the nursing team and the transfusion medicine laboratory to ensure no gaps in release of laboratory test results or transfusion support.				
40. Tranexamic acid should be administered as soon as intravenous or intraosseous access is achieved but within 3 hours from time of injury or within 3 hours from MHP activation in all other patients.	Tranexamic acid reduces mortality in the setting of trauma ⁹³ and postpartum hemorrhage. ⁹⁴ It is most effective when given immediately, with the survival benefit decreasing by 10% for every 15-minute delay in administration and with no benefit after 3 hours from injury/onset of bleeding. ⁹⁵ There is no increased risk of venous or arterial thromboembolic complications. ⁹⁶ Dosages and infusion rates vary depending on the study protocol (1-g bolus plus 1-g infusion over 8 h, ⁹³ 1-g bolus and 1-g bolus repeated at 1 h, ⁹⁷ 1-g bolus and repeated if ongoing bleeding at ≥ 30 min ⁹⁴ and 2-g bolus at the scene of the injury (Clinicaltrials.gov; NCT01990768; trial completed). The dosage and infusion rate should be determined by the local institution. Simplification may be needed in more resource challenged locations, and a single 2-g bolus may be preferred. Evidence of the efficacy of tranexamic acid in pediatric trauma is currently limited, but its use in pediatric patients with trauma requiring transfusion is accepted practice, within the same time parameters as for adults. For pediatric patients, the initial bolus of tranexamic acid can be dosed at 15 mg/kg up to a maximum of 1 g, with additional doses/infusion based on local policy. ^{96,99} The role of tranexamic acid in gastrointestinal bleeding has not been confirmed; a large multicentre trial (Haemorrhage Alleviation With Tranexamic Acid – Intestinal System [HALT-IT]) is under way to determine whether tranexamic acid assists with hemostasis and reduces transfusion or mortality rates. ¹⁰⁰ Tranexamic acid should be readily available in clinical areas where massive hemorrhage is common to prevent delays in administration.				
41. MHP activations should be reviewed by a multidisciplinary committee for quality assurance.	Compliance with MHPs is poor during the resuscitation of a critically ill patient who has multiple competing priorities. ^{12,16} Implementation of an MHP is just the first step to improving the care of massively bleeding patients; training, simulations, checklists, audit and feedback are needed to achieve high levels of performance. At a minimum, the quality metrics listed in Box 1 should be tracked on consecutive MHP activations by a multidisciplinary team, with feedback to the front-line staff at regular intervals.				
42. The quality metrics in Box 1 should be tracked on all activations of the protocol and the data reviewed quarterly at the hospital transfusion committee and the medical advisory committee.	_				
Note: aPTT = activated partial thromboplasti RBC = red blood cell.	n time, INR = international normalized ratio, MHP = massive hemorrhage protocol, PCC = prothrombin complex concentrate,				

*Few patients will require more than 12 units of RBCs for an acute hemorrhage. In patients who have received 12 or more units of RBCs, transfusion decisions for plasma and fibrinogen replacement should be made based on hourly measurement of the INR and fibrinogen levels, and orders communicated promptly to the blood bank

Interpretation

Through a modified Delphi iterative process, we selected and refined 42 statements and 8 quality indicators to form the foundation for the proposed provincial standardized, evidence-based MHP template for hospitals. The MHP toolkit will include training material, simulation exercises, checklists, template policies and procedures, and patient material. In addition to decreasing variability in care, reducing cognitive load on providers, improving communication between the clinical and laboratory teams, increasing uptake of evidence-based treatments and ultimately improving patient outcomes, we hope that this initiative will improve patient comfort and safety, communication with families and disclosure of transfusion risks to patients. Although the lists of statements are not exhaustive, we hope that they address the current widespread variability in MHP structure. 14,15 Both the modified Delphi process and the external stakeholder consultation assisted with statement construction to ensure clarity for both experts in transfusion medicine and health care personnel working outside of the laboratory.

This consensus document created by the modified Delphi method is a powerful tool that is broadly applicable and adaptable to many hospital settings. The consensus panel included excellent representation by resource-limited and geographically constrained sites in rural Ontario, as well as resource-rich





Box 1: Quality metrics to be tracked on all activations of the protocol				
Quality metric	Local reporting	Provincial reporting		
Q1. Proportion of patients receiving tranexamic acid within 1 h of protocol activation	X	Х		
Q2. Proportion of patients in whom RBC transfusion is initiated within 15 min of protocol activation	Х	Х		
Q3. Proportion of patients (of patients requiring transfer for definitive care) with initiation of call for transfer within 60 min of protocol activation	X			
Q4. Proportion of patients achieving temperature ≥ 35°C at termination of the protocol	Х			
Q5. Proportion of patients with hemoglobin levels maintained between 60 and 110 g/L during protocol activation, excluding certain pediatric populations (e.g., neonates) that may require higher hemoglobin values	Х			
Q6. Proportion of patients transitioned to group-specific red blood cells and plasma within 90 min of arrival/onset of hemorrhage	Х	Х		
Q7. Proportion of patients with appropriate activation (≥ 6 RBC units in first 24 h, > 40 mL/kg per 24 h of RBCs in pediatric patients) or before this level in patients dying due to hemorrhage within 24 h	Х			
Q8. Proportion of patients without any blood component wastage (including plasma that is thawed and not used within the 5-day limit on another patient)	Х			

tertiary care centres. As a result, the recommendations reflect best practice but are also flexible, as they include modifiers for clinical environments with fewer team members and limited access to laboratory testing and blood product availability. For example, the MHP recommends that centres without the ability for definitive hemorrhage control call for early transport (statements 1 and 10), facilities where certain blood components are limited are offered alternatives (statement 34), and there is dosage modification for the initial antifibrinolytic agent for patients in remote locations to ensure administration of a timely and complete dose (statement 40). In comparison, a recent survey of hospitals with MHPs in the United States showed that the majority were in large academic centres, calling for a higher ratio of plasma:red blood cells (1:1), with very few unified MHPs in place to encompass nontrauma indications. ¹⁰¹

The Delphi technique was chosen as the method to develop consensus statements from expert stakeholders because of the absence of clinical trial evidence for all aspects of the management of the massively bleeding patient. Although clinical trials provide considerable guidance on the utility of blood component ratios, antifibrinolytic agents, as of recombinant factor VIIa and other areas of management, they fail to provide recommendations on how to construct the protocol, modifications for community hospitals or specific patient populations. Although the Delphi exercise to allow for an open forum after round 1 to increase input from experienced practitioners and to discuss the vast quantity of available literature to ensure that all participants had a foundation in MHP knowledge. This allowed for the

broadest capture of areas of massive hemorrhage that are logistical in nature and do not lend themselves to evaluation in clinical trials, such as communication strategies and procedures, protocol nomenclature, frequency and type of laboratory testing, laboratory resuscitation targets, and blood component transport and bedside storage. Importantly, modification of the Delphi exercise also allowed the invaluable input of a patient representative, who provided insight into the importance of patient-related outcomes such as communication and hypothermia management to ensure comfort.

Limitations

The major limitations and challenges of our modified Delphi process included the limited number of panellists per specialty and hospital type (e.g., obstetrics, remote hospitals) owing to cost and logistical limitations. Furthermore, 1 panellist ranked the statements based on feasibility and/or cost of recommendations despite instructions to rank based on best practices; however, later in round 1, the panellist reranked based on best practices. Another limitation was the failure to include panellists involved in hospital blood/sample transportation and communications, as these colleagues are critically important to the success of the MHP (e.g., ensuring a constant supply of blood packs from the bank). Given the numerous items lacking clinical trial evidence, our recommendations will need to be revisited and updated at regular intervals to evolve with this rapidly changing field of medicine. We note a particular lack of high-quality evidence for pediatric patients owing to exclusion of these patients from many clinical trials.



Conclusion

We are confident that the 42 statements and 8 quality indicators, constructed through a highly structured process and with the involvement of a diverse and knowledgeable group of experts, will serve as a strong foundation for the creation of a robust MHP toolkit. The MHP toolkit, currently under development, will consist of recommendations and educational resources for 12 sections (patient transport; principals of damage control resuscitation; team dynamics; communication; laboratory testing; temperature management; use of blood components and adjuncts, including anticoagulant reversal; simulation and training tools; pediatric-specific recommendations; patient communication follow-up tools; research goals; and quality metrics and reporting) written by an expert working group, to be published and freely accessible in the spring of 2020 on the Ontario Regional Blood Coordinating Network website (http://transfusionontario.org/en/) and presented at the network's spring Transfusion Committee Forum. We expect that, with the use of the toolkit, hospitals will achieve higher adoption of evidence-based care of the patient with massive hemorrhage, improved speed of delivery of blood components and hemostatic adjuncts, and more diligent monitoring of clinical and laboratory parameters. There is an opportunity to track patient outcomes in existing prospectively collected databases in trauma and obstetrics and for inpatients held by the Canadian Institutes for Health Information in Ontario to understand the impact of this effort to standardize the care of these complex, high-acuity patients.

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