Assessing and Replenishing Lost Volume

J. G. L. Cockings and C. S. Waldmann

INTRODUCTION

Classically, shock is defined as a state of inadequate tissue perfusion in relation to the metabolic needs of a given patient. Inadequate blood flow may manifest clinically as tachycardia, pallor, oliguria, altered mental status, the development of lactic acidosis, or a combination of these changes.

From a physiological point of view, shock is hypovolemic, cardiogenic, anaphylactic or cytotoxic. Hypovolemic shock is classically associated with postpartum hemorrhage (PPH) and due to loss of circulating blood volume. Hypotension is often present in severe cases, but is a late sign and is a poor guide to the volume of blood lost, as pregnancy is accompanied by an alteration of cardiovascular physiology, and the response to blood loss and its management may differ from the non-pregnant situation.

In the UK today, massive PPH accounts for 35% of obstetric admissions to intensive care1–3. The 2006–2008 Confidential Enquiry into Maternal Deaths1 shows that PPH continues to be a significant cause of peripartum maternal deaths, despite the relative luxury of equipment, staffing and other resources. It is axiomatic that bleeding patients demand rapid assessment and judicious replenishment of lost circulating volume, albeit within the context of the compensatory effects of hypovolemic shock and the physiological changes that occur in late pregnancy.

PHYSIOLOGY

The normal circulating blood volume for a healthy non-pregnant adult is 70 ml/kg, or 7.5% of body weight. Cardiac output is 4–6 l/min, and the non-pregnant adult systemic vascular resistance is 10–15 mmHg/l/min (900–1200 dyne.s/cm5). Maternal blood volume increases during pregnancy to 40% above baseline by the 30th week, with an accompanying but smaller (20–30%) increase in red cell volume. Cardiac output increases to 50% above pre-pregnancy levels by the 24th week. Systemic blood pressure is more variable in healthy uncomplicated pregnancy, with a small fall in the first and second trimesters, but return to pre-pregnancy levels by the third. Resting heart rate increases progressively in the first and second trimesters to 15–20 bpm above pre-pregnant levels. In addition to these changes, others take place in the autoregulation of intravascular volume and the circulation, both of which affect the body’s response to blood loss. Examples include a blunted response to angiotensin II, which in part may be due to an increased production of nitric oxide4, a decreased tolerance to postural changes and an increased cardiac noradrenaline turnover5,6.

Circulating volume, clinical signs of hypovolemia and the body’s ability to compensate for volume loss are also all affected by pregnancy related diseases and their treatment, the effects of which continue on into the early postpartum period. Pre-eclampsia, for example, causes a contracted effective arterial blood volume compared with the normal peripartum state. Vascular reactivity is increased, and widely used drugs such as hydralazine and magnesium compromise the body’s ability to produce compensatory vasoconstriction in the face of hemorrhage. Indeed, it appears that there is a failure to increase plasma volume and reduce systemic vascular resistance in pre-eclampsia due to inadequate trophoblastic invasion into the spiral arteries of the uterus6. Pre-eclamptic patients thus have an increased tendency to develop pulmonary edema during volume replacement due to many factors, including increased capillary permeability, hypoalbuminemia and left ventricular dysfunction6.

Normal delivery results in predictable blood losses which range from 300–500 ml for vaginal deliveries to 750–1000 ml for cesarean section births, although these numbers are variously described not only in this text, but also in the literature published in the past 5 years (see Chapter 11). Regardless, in addition to blood lost from the body, a substantial amount of blood is also redirected into the systemic circulation, often referred to as the autotransfusion effect. This results in an increase in cardiac output by as much as 80% (see Chapter 22). The effect persists in uncomplicated patients, gradually returning to non-pregnant levels at 2–3 weeks6.

ASSESSMENT OF CIRCULATING BLOOD VOLUME

Young healthy adults can compensate for the loss of large volumes from the circulation with few initially obvious external signs. To say that accurate assessment of blood loss is difficult for the experienced as well as the inexperienced examiner is not only correct but...
also represents an important theme that is re-echoed in numerous chapters of this book.

In cases of hemorrhage, symptoms often precede signs. These include unexplained anxiety and restlessness, the feeling of breathlessness (with or without an increased respiratory rate), and a sensation of being cold or generally unwell. For healthy, non-pregnant adults, hypovolemia and associated signs can be divided into four stages (Table 1). These range from the largely undetectable stage 1 with less than 15% loss of volume, to the severe life-threatening stage 4 when more than 40% has been lost. Unfortunately, comparable tables for early and late pregnancy and the immediate postpartum period have not been compiled, but the signs follow a similar pattern.

As helpful as such tables may be, the most important clinical principle in the treatment of PPH is early recognition and prompt correction of lost circulating volume, together with simultaneous medical and/or surgical intervention to prevent further loss. Early recognition of life-threatening physiological derangements can be improved by the use of early-warning scoring systems.

Recording physiological observations at regular intervals has long been routine practice in hospitals. Early-warning scores derived from simple routine physiological recordings can identify those patients with greater risk of critical illness and mortality. In recognition of the normal physiological changes in pregnancy, specialized scores have been developed for use in the obstetric population. These Modified Early Obstetric Warning Scores (MEOWS) are increasingly used in obstetric units throughout the UK and other jurisdictions. Their real value is that they can flag the early but sometimes subtle signs of concealed and largely compensated hemorrhage in the early postpartum patient. These scores use the physiological parameters most likely to detect impending life-threatening compromise. They are based on simple physiological observations which do not demand special skills, thus allowing them to be used across all health care systems, rich or poor. The variables usually comprise respiratory rate, heart rate, systolic blood pressure, temperature and mental awareness, each being assigned a weighted score while the total score is the sum of these. Such systems are reproducible and effective at predicting the likelihood of progression to critical illness. They also are well suited to the early detection of the often subtle signs of unappreciated blood loss. Their use allows a trigger value for ward staff to call for assistance from intensive care or other senior staff. An example of an early warning score for the obstetric population used in the UK is given in Table 2.

Once the possibility of intravascular depletion has been raised, a prompt clinical assessment is urgent, as the clinical condition of the patient can change rapidly. Clinical assessment, in association with non-invasive and invasive monitoring where appropriate, must be made by senior clinicians (if available), with special attention to repeated assessment at frequent intervals to detect the problem as early as possible. If senior clinicians are not available, they should be notified as described in the protocols in Chapters 40 and 41.

Clinical examination is performed simultaneously with incident-related history taking. This history may elicit obvious features associated with shock such as overt blood loss and pain, but may also elicit more subtle features such as general malaise, anxiety and restlessness, a poorly defined sense of doom and breathlessness. Physical examination is directed to the fundamental areas of vital function, the conscious state and airway protection, the adequacy of respiratory function, oxygenation and circulation. In particular, the following should be assessed and documented:

1. Early stages of shock are associated with restlessness and agitation, sometimes with a heightened sense of thirst, but these progress to drowsiness when around 30% of blood volume is lost. Loss of consciousness is a very late sign, with significant risk of imminent death.

2. Tachypnea is an early sign, partly driven initially by the anxiety, but is an independent sign, and the respiratory rate increases with progressive blood loss and will usually exceed 20 breaths/min when 30% of blood volume is lost.

3. Oxygenation becomes harder to assess clinically as peripheral pallor becomes more marked, and the pulse oximeter becomes less reliable as peripheral perfusion becomes weaker.

4. A fall in the jugular venous pressure occurs reasonably early, but is partly compensated for by a

<table>
<thead>
<tr>
<th>Classification</th>
<th>Stage 1</th>
<th>Stage 2</th>
<th>Stage 3</th>
<th>Stage 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood loss (% volume lost)</td>
<td>10–15</td>
<td>15–30</td>
<td>30–40</td>
<td>&gt;40</td>
</tr>
<tr>
<td>Conscious state</td>
<td>Alert, mild thirst</td>
<td>Anxious and restless</td>
<td>Agitated or confused</td>
<td>Drowsy, confused or unconscious</td>
</tr>
<tr>
<td>Respiratory rate</td>
<td>Normal</td>
<td>Mildly elevated</td>
<td>Raised</td>
<td>Raised</td>
</tr>
<tr>
<td>Complexion</td>
<td>Normal</td>
<td>Pale</td>
<td>Pale</td>
<td>Marked pallor or gray</td>
</tr>
<tr>
<td>Extremities</td>
<td>Normal</td>
<td>Cool</td>
<td>Pale and cool</td>
<td>Cold</td>
</tr>
<tr>
<td>Capillary refill</td>
<td>Normal</td>
<td>Slow (&gt;2 s)</td>
<td>Slow (&gt;2 s)</td>
<td>Minimal or absent</td>
</tr>
<tr>
<td>Pulse rate</td>
<td>Normal</td>
<td>Normal</td>
<td>Elevated</td>
<td>Fast but thready</td>
</tr>
<tr>
<td>Systolic blood pressure</td>
<td>Normal</td>
<td>Normal</td>
<td>Normal or slightly low</td>
<td>Hypotensive</td>
</tr>
<tr>
<td>Urine output</td>
<td>Normal</td>
<td>Reduced</td>
<td>Reduced</td>
<td>Oliguric</td>
</tr>
</tbody>
</table>

Modified from Baskett, 1990.

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Modified early obstetric warning score. Reproduced with permission by Dr. R. Jones, Consultant Anaesthetist, Royal Berkshire Hospital, UK.

Table 2  Modified early obstetric warning score. Reproduced with permission by Dr. R. Jones, Consultant Anaesthetist, Royal Berkshire Hospital, UK

<table>
<thead>
<tr>
<th>Score</th>
<th>3</th>
<th>2</th>
<th>1</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiratory rate (bpm)</td>
<td>&lt;8</td>
<td>9–18</td>
<td>19–25</td>
<td>26–30</td>
<td>&gt;30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pulse rate (bpm)</td>
<td>&lt;40</td>
<td>40–50</td>
<td>51–100</td>
<td>101–110</td>
<td>111–129</td>
<td>&gt;129</td>
<td></td>
</tr>
<tr>
<td>Systolic blood pressure (mmHg)</td>
<td>&lt;70</td>
<td>71–80</td>
<td>81–100</td>
<td>101–164</td>
<td>165–200</td>
<td>&gt;200</td>
<td></td>
</tr>
<tr>
<td>Diastolic blood pressure (mmHg)</td>
<td>&lt;95</td>
<td>95–104</td>
<td>&gt;105</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conscious level</td>
<td>Unresponsive</td>
<td>Responds to pain</td>
<td>Responds to voice</td>
<td>Alert</td>
<td>Irritated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Urine hourly (ml/h) or in 24 h</td>
<td>&lt;=30</td>
<td>&lt;45</td>
<td>&gt;45</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(&lt;720 ml)</td>
<td>(&lt;1000 ml)</td>
<td>(&gt;1000 ml)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Final score = sum of individual scores at any one time

Action:
Score 0 or 1 Repeat observations when appropriate for clinical scenario
Score 2 Inform midwife in charge, repeat in 15 min
Score 3 Inform midwife in charge, obstetric registrar and duty anesthetist
Score 4 As above but the consultant obstetrician should be informed
Consider informing duty consultant anesthetist and intensive care team

*The timing of involvement of the most senior obstetrician and anesthetist, as well as the timing of intensive care involvement will be highly dependent on the staffing structure, seniority and operating principles in the particular institution. Each institution, if introducing an Early Warning Score system, should define the callout thresholds appropriate for their own particular organizational structure.

Assessing and Replenishing Lost Volume

MANAGEMENT

When the compensating mechanisms directed toward maintaining the blood pressure are exhausted, pressure readings fall dramatically. At this point, shock is advanced and the risk of imminent death is significant. If the patient is under competent medical care, however, by this same point in time, the significant blood loss will have been recognized, volume and, in the majority of instances, blood replacement begun via large-bore peripheral access and medical therapies and surgical intervention organized. Clinical signs alone, even by experienced clinicians, are often unreliable in estimating the volume deficit and the degree of shock when well compensated. Therefore, other methods to assess more accurately volume status and circulatory adequacy must be used to aid clinical assessment.

The first and simplest of these is invasive measurement of central venous pressure. A central venous catheter can be placed in any central vein, but it should be remembered that, once the patient is in actual hypovolemia, identification of a central vein may be difficult without the use of ultrasound. The internal jugular vein is the preferred site in this situation, as the femoral vein is relatively inaccessible, and the subclavian route may have a higher risk of complications in late pregnancy and the early postpartum period, especially if inserted under urgent conditions.

The National Institute for Clinical Excellence (NICE) suggests that cannulation of central veins using two-dimensional ultrasound imaging is the preferred method, with the evidence strongest for the internal jugular route. In the healthy, non-pregnant adult, the systemic venous capacity is 3–4 liters, or 75% of circulating volume. If the tone of the venous capacitance vessels did not change as volume was lost from the circulation, the central venous pressure would fall quickly and early, with early compromise of the cardiac output. However, as blood volume is lost, the tone in these venous capacitance vessels increases, moving blood centrally, and maintaining central venous pressure.

Confusion surrounding the concept of venous return can be dispelled if it is thought of in terms of right atrial pressure rather than an increased flow of blood to the right atrium. As blood is lost, the volume in the venous capacitance vessels is reduced and the tone in these vessels increases. The central venous pressure falls progressively, but to a lesser degree due to the compensatory increase in this venous tone. Figure 1 shows the relationship between venous capacitance and central venous pressure during acute blood loss and immediate replacement. As blood is lost from the circulation, the patient follows the line A to B (Figure 1). The central venous pressure falls slowly at first, then more steeply as the extent of blood loss increases. As volume is returned to the circulation, the patient will follow first the line B to C and then C to D, rather than simply returning from B to A. This is
Hemoglobin estimates indicate the concentration of hemoglobin in the sample. The simplicity of this statement only reinforces the point that the concentration of hemoglobin following acute blood loss reflects either medical intervention or the patient’s compensation to blood loss. Acute blood loss alone will not change the concentration of hemoglobin in the blood left in the system. The concentration is reduced only when the lost volume is replaced by internal fluid shifts or external fluids are added to the system that are low in hemoglobin. If left untreated, however, acute blood loss results in a fall in hemoglobin concentration after about 4–6 h due to internal compensatory fluid shifts. In contrast, intravenous administration of fluid will dilute the hemoglobin more quickly.

Further information regarding the status of the circulation can be gained by more invasive or complex techniques. These include the pulmonary artery catheter, pulse contour analysis and the esophageal Doppler technique. The measurement of cardiac output by means of the thermodilution technique using a pulmonary artery catheter was first described by Bradley and Branthwaite in 1968 and subsequently popularized by Swann, Ganz and co-workers. Despite being the gold standard for invasive assessment of cardiac output and left atrial pressure for several decades, some evidence suggests that the risks of this technique may outweigh its benefits in many circumstances. The benefit of the pulmonary artery catheter is that it provides reliable measurements even in the face of changing body position and is equally effective in the awake, conscious patient as well as the sedated or anesthetized patient receiving artificial ventilation. The risks include those associated with the insertion of a large-bore cannula into a central vein, which is higher risk in the volume-depleted patient suffering massive blood loss, even with the aid of the two-dimensional ultrasound technique. Risks also relate to infection, cardiac arrhythmias, pulmonary artery damage and lung injury. To date, experience with the pulmonary artery catheter in the obstetric population has been in patients with pre-eclampsia and eclampsia rather than massive PPH.

Fortunately, pulse contour analysis is a realistic alternative to the pulmonary artery catheter. This system requires only standard peripheral arterial and central venous cannulae. A common example of this technique uses cardiac output estimated initially and periodically thereafter by the lithium dilution technique (LiDCO Ltd., Cambridge, UK). The pressure waveform is analysed using this static cardiac output measurement as a reference to estimate a stroke volume. Changes to the pulse waveform and heart rate from this point are used to estimate stroke volume and cardiac output on a continuous display. Calibration is performed by periodic re-assessment of the cardiac output using the lithium dilution technique. Limitations relate to issues of non-linearity or aortic compliance, how closely the radial arterial pulse waveform resonance relates to the proximal aortic waveform and, therefore, stroke volume, the common problems of
damped arterial waveforms, drift between static measurements of cardiac output, and problems associated with poor transmission of pulse waves in severe arrhythmias. Despite these concerns, this technique can provide a continuous idea of cardiac output, systemic blood pressure and vascular resistance in any patient with central venous and peripheral arterial access and is well suited for monitoring the postpartum patient who has undergone massive hemorrhage and is undergoing resuscitation. Other commercially available pulse contour analysis systems use alternative methods for measurement of the reference cardiac output. An example is the PiCCO system (Pulsion Medical Systems, Munich, Germany) which employs a transpulmonary thermodilution measurement from an axillary or femoral artery.

**REPLENISHING LOST VOLUME**

[Editor's Note: This topic is under intense debate on an international basis. A thorough discussion is provided by Lockhart in Chapter 4 and Paidas in Chapter 6. L.G.K.]

Replacing lost circulating volume should commence as soon as significant bleeding is recognized and, ideally, before the signs of significant hypovolemia have developed. Important initial measures are to simultaneously provide supplemental oxygen, ensure multiple large-bore peripheral intravenous access, undertake an initial rapid clinical assessment and summon senior members from anesthesia, intensive care, surgical and hematology departments for assistance when these individuals are available. Each institution should have a rapid response protocol in place for the management of massive hemorrhage and PPH in particular. This protocol should be familiar to all, easily accessible and followed (see Chapters 36 and 40).

The principal underlying aim of volume replacement during and following massive PPH is restoration and maintenance of tissue perfusion to all body organs in order to maintain cellular function and viability. Although the initial focus should be on restoration of the common clinical indicators of shock, the clinician must proceed further. Even if all conventionally used criteria resolve, shock may still be present on a cellular, the common clinical indicators of shock, the clinician

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Initial volume replacement enhances right atrial filling and improves cardiac output. As shock develops with blood loss, venous tone increases as described above. Volume administration should be rapid, but titrated to the right atrial filling pressure. Initially, right atrial filling pressure may be restored by a smaller volume than that lost due to the reduced capacity in the venous capacitance vessels. Indeed, immediate rapid re-infusion of the entire volume lost may provoke fluid overload if the tone in the capacitance vessels did not decrease as rapidly. This can be appreciated from Figure 1; here, rapid volume replacement occurs along the line B to C. Central venous pressure rises despite the intravascular compartment remaining depleted. As the venous tone relaxes following resuscitation, further volume administration can occur with a central venous pressure falling toward normal as the volume in the venous capacitance becomes replete (line C to D). Thus, volume administration should be rapid but infused in discrete volume challenges, with the effect on the right atrial filling pressure, systemic blood pressure and other hemodynamic variables being monitored. Commonly, 250–500 ml of either a crystalloid or a colloid is administered over a period of 10–20 min as the urgency dictates and while blood is being obtained (a patient with life-threatening stage 4 shock will receive 2–3 liters more quickly, but, even then, the principles of monitoring the hemodynamic variables during the infusion of fluid remain). Simple measures of tissue underperfusion, which may persist after apparent restoration of global hemodynamics, include the base deficit and serum lactate. Efforts to measure and enhance tissue perfusion should continue until all such parameters return to normal. More specific measures to monitor tissue perfusion, including tissue oxygen tension devices and gastric tonometry, are not widely used.

The best fluid to use for volume expansion in hemorrhagic shock remains a matter of debate. Both crystalloid and colloid are effective, but each has advantages and disadvantages (Table 3).

One recent large study showed no difference in mortality in intensive care patients requiring volume expansion whether this expansion was made with saline or albumin. Colloids expand the intravascular space preferentially, whereas crystalloids quickly become distributed throughout the extracellular space. Saline has the disadvantage of hyperchloremia, which causes a dilutional or hyperchloremic acidosis. The use of crystalloids is not associated with anaphylaxis, whereas colloids such as the gelatins can produce severe life-threatening reactions, although this is less common with hydroxyethyl starch. Crystalloids have minimal effect on coagulation other than a dilutional effect, although saline infusions may have a procoagulant effect. Overall, crystalloids have a lower cost and lower incidence of side-effects, but the colloids have several theoretical advantages regarding tissue edema and oxygen delivery to the tissues. Despite intense debate and research interest, neither crystalloids nor colloids have been shown to be...
superior to one another regarding survival outcome from hemorrhagic shock.

Regardless of which substance is selected for volume expansion at the start of therapy, it is essential that blood be administered and that a protocol be available for the use of blood products in instances of massive bleeding. In the UK, the responsibility for maintaining such a protocol lies with the hospital blood transfusion committee, a multidisciplinary committee that all hospitals must by law ensure is in place and answerable to the hospital executive. It is unacceptable to have situations where the laboratory insists on blood samples being sent for blood count and coagulation studies before any blood products are issued; the on-call hematology consultant should be actively involved and aid with the use of blood, fresh frozen plasma, platelets, cryoprecipitate and the use of recombinant activated factor VII.

The Trendelenburg position is often used in the management of the hypotensive patient, but its benefit has been questioned. The concept is to displace blood from the lower limbs centrally, to increase preload and enhance cardiac output as a temporary measure until adequate blood volume can be restored. However, little proof exists that this theoretical benefit occurs in practice. Sibbald and colleagues in 1979 showed that, in hypotensive patients, the Trendelenburg position did not significantly increase preload, but did increase afterload and blood pressure at the expense of cardiac output. A review of available data concludes that the Trendelenburg position ‘is probably not a good position for resuscitation of patients who are hypotensive’.

The conventional approach to severe hemorrhage, where the endpoint is euvoemlia with restoration of a normal blood pressure, heart rate and cardiac output, has been questioned in the out-of-hospital trauma setting. Although not based on evidence from the obstetric population, the physiological rationale may still be applicable. Falling blood pressure and cardiac output, together with increased sympathetic tone and release of endogenous catecholamines, reduce the rate of blood loss. Restoration of these parameters without control of the bleeding will increase the total volume of blood loss, increasing the degree of coagulopathy, reducing oxygen-carrying capacity and ensuing multi-organ dysfunction. Low-volume fluid resuscitation for hemorrhagic shock may be a possibility and the evidence suggests that volume resuscitation should be deliberately limited to the minimum required to sustain vital organ function until the bleeding has been arrested, such as by surgery.

Small-volume hypertonic resuscitation also has been advocated for hemorrhagic shock. Here too, the target population was not obstetric, and the shock was not from PPH. The concept is that a relatively small infused volume will cause much larger expansion of the circulation by drawing water into the intravascular compartment. There is evidence that there may be beneficial effects of endothelial and red cell edema and capillary flow, but concerns are present regarding other potentially adverse effects such as that on the immune system. This latter concern has not been shown to be a problem in clinical practice.

Maintenance of the hemoglobin concentration is essential for oxygen-carrying capacity and delivery to the tissues. Titration of fluid and blood products to an exact hemoglobin level in a rapidly bleeding patient is difficult. A hemoglobin level of 7–8 g/dl appears an appropriate threshold for transfusion in the intensive-care population, with possible benefit for a higher level of 9 g/dl for those with ischemic heart disease. It is logical to aim at the high end of the target range when resuscitating from hemorrhagic shock as there is a tendency to drift down. A target of 10 g/dl has been suggested as a reasonable goal in the actively bleeding patient. In the case of PPH, however, the goal is two-fold cessation of the hemorrhage as well as restoration of the hemoglobin level.

Coagulation disorders are both predisposing factors for, and consequences of, massive PPH. A bleeding diathesis from a coagulopathy, thrombocytopenia or platelet dysfunction may result from pre-existing disease, a pregnancy-acquired disorder, such as eclampsia, or treatment, such as aspirin. Massive blood loss also

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**Table 3 Intravenous fluids**

<table>
<thead>
<tr>
<th>Type of fluid</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crystalloids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Saline</td>
<td>Cheap; easily available; long history of use</td>
<td>Produces a hyperchloremic acidosis; small procoagulant effect</td>
</tr>
<tr>
<td>Hartmann’s</td>
<td>No risk of anaphylaxis; minimal direct effect on the base deficit; easily available</td>
<td>Mildly hypotonic</td>
</tr>
<tr>
<td>5% Dextrose</td>
<td>No place in acute expansion of the intravascular space</td>
<td>Hypotonic; no significant expansion of the vascular space; rapid distribution to intracellular and extracellular spaces</td>
</tr>
<tr>
<td>Hypertonic saline</td>
<td>Rapid expansion of the intravascular space in excess of the volume infused; possible beneficial effects on red cell and endothelial edema and capillary blood flow</td>
<td>Insufficient data; uncertainty regarding possible adverse effects such as on the immune system</td>
</tr>
<tr>
<td>Colloids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gelatins</td>
<td>Largely remains in the intravascular space for 2–4 h</td>
<td>Risk of anaphylaxis; no clear survival advantage over crystalloids</td>
</tr>
<tr>
<td>4% Human albumin</td>
<td>More physiological than gelatins; remains predominantly in the intravascular space for 12 h</td>
<td>Expensive; no clear survival advantage over crystalloids</td>
</tr>
<tr>
<td>Hydroxethyl starch</td>
<td>Remains in the intravascular space for 12–24 h</td>
<td>Risk of coagulopathy, renal injury and reticulo-endothelial accumulation</td>
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</tbody>
</table>
creates both a coagulopathy and thrombocytopenia through dilution and consumption. These issues and their management are discussed in detail in Chapters 25 and 50.

SUMMARY

Rapid assessment of the presence of occult bleeding or intravascular volume depletion is essential. The body can compensate for blood loss such that, by the time obvious clinical signs are present, a significant volume can already be lost and tissues already in a state of hypoperfusion. Normal physiological adaptations in late pregnancy that persist into the postpartum period can make recognition and quantification of intravascular loss difficult, and can render the body less capable of withstanding massive blood loss. This can be further complicated by pregnancy-related disease such as pre-eclampsia and its treatment, and modalities such as hydralazine and magnesium.

Assessment of both the degree of loss and the response to volume replacement require clinical skills, invasive hemodynamic monitoring and the early involvement of senior clinicians. The use of Modified Obstetric Early Warning Scores in all patients to aid the early detection of concealed hemorrhage and serious acute illness is to be strongly encouraged. A simple system such as this based on easy to measure physiological variables is easy to implement, does not rely on extensive training or experience and does not need to be limited to the developed world.

There is no one correct fluid to use. It is usual to use a combination of crystalloids or colloids and blood products to maintain a hemoglobin concentration of near 10 g/dl during the actively bleeding period (7–9 g/dl is probably safe once the active bleeding has been stopped). Coagulopathies and thrombocytopenia also need to be corrected with appropriate transfusion products and with active involvement of the hematologists. There may be a place for limited volume expansion before the bleeding has been stopped surgically to reduce the volume lost, but this must not be at the cost of demonstrable organ ischemia.

Prompt recognition, close monitoring of volume status, rapid arrest of the bleeding and adequate volume resuscitation are all required but when used together can reduce mortality from PPH.

References

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