Several aspects of the management of an orthopaedic surgical patient are not directly related to the surgical technique but are nevertheless essential for a successful outcome. Blood management is one of these. This paper considers the various strategies available for the management of blood loss in patients undergoing orthopaedic and trauma surgery.

Blood management is now receiving more attention, as shown by the increasing number of relevant publications over the past few years.1-5 There has thus been increased awareness not only of the amount of blood actually lost in common orthopaedic procedures but also of the possible disadvantages of allogenic blood transfusion, resulting in a general trend towards avoiding or minimising its use.

Potential risks associated with blood transfusion

Historically, correction of blood loss in surgical patients has been by transfusion of allogenic blood to maintain or to restore the haematocrit and level of haemoglobin (Hb) close to normal. It is now accepted that the historical trigger values of 30% haematocrit and 10 g/dl of Hb can safely be lowered to 8 g/dl and even to 7 g/dl of Hb in a patient with no major comorbidities.6 Experience with surgery on patients who are Jehovah’s Witnesses has shown that mortality only becomes a major risk when the pre-operative Hb level is under 6 g/dl7,8 and also that elective surgery such as total hip (THR) or knee replacement (TKR) can be performed safely without blood transfusion in non anaemic patients who refuse blood transfusion.9

Other reasons contributing to the decline in the use of transfusion of allogenic blood include the effect on the immunological system, with various possible adverse consequences as follows.

A correlation has been reported between allogenic blood transfusion and post-operative infection in the elective surgical patient10-14 and even more so in the traumatically-injured patient.11,13 Leucocyte reduction by filtration has been found to reduce this effect without abolishing it.14 A number of countries have now opted for the general usage of leucocyte-depleted blood in surgery. The association between the transfusion of non-leukodepleted allogenic blood and infection is being questioned, however, since its demonstration may have been biased for methodological reasons.15 Increased recurrence rates in patients with malignant tumours have been related to immunomodulation resulting from allogenic blood transfusion16,17 and other effects such as multiple-organ failure or mortality have been attributed to immunomodulation from allogenic blood transfusion by pro-inflammatory mechanisms.16-18

There is a risk of viral transmission of HIV, hepatitis C virus, and hepatitis B virus although this has been dramatically reduced by the screening of donors and the development of efficient serological tests. In 2005, the risk was estimated at 0.31 per million blood donations for HIV, 0.10 per million donations for hepatitis C virus and 1.44 per million donations for hepatitis B virus in France,19 with similar figures in the United Kingdom of 0.14, 0.80 and 1.66 per million donations for HIV, hepatitis C virus and hepatitis B virus, respectively.20 The possibility of the transmission of prions by allogenic blood transfusion has now been established in a small number of cases.21 The risk of bacterial contamination of blood products appears to affect platelet concentrates (1/50 000) much more than leukocyte-depleted erythrocyte concentrates (1/500 000).22

Administrative errors in the handling of donated blood are rare but are occasionally responsible for major incidents which may be lethal. The risk applies to autologous blood transfusion also. In an evaluation of near-miss events in 35 French hospitals, Chironi et al23...
found that in 407 147 compatibility tests the incidence of ABO incompatibility was one in 3400, which was related to phlebotomy errors or to identification errors, but the incidence of ABO mismatched transfusion was ten times lower.

Allogenic blood transfusion also adds to the cost of an operation, all the more as it has been shown to be associated with increased morbidity and prolonged hospital stay. 24

**Elements in the reduction of need for allogenic blood transfusion**

A number of clinical studies 1-5 have investigated ways of reducing the need for allogenic blood transfusion after orthopaedic operations while minimising post-operative anaemia. Most of these have focused on one specific action, such as pre-operative autologous blood donation, intra-operative cell saving, or re-infusion of drained blood. Blood management is a complex issue, however, and an integrated approach is necessary to address all its components.

Allogenic blood transfusion is required when a patient has lost more red blood cells than his/her condition would allow without endangering life. It is unavoidable in a number of patients after multiple trauma involving, for instance, complex fractures of the pelvis. In elective surgery, the risk can be evaluated well in advance, and a number of measures can be considered in order to reduce the probability of having to resort to allogenic blood transfusion. The key elements which should be taken into consideration are the pre-operative erythrocyte stock, the anticipated peri-operative blood loss and the acceptable blood loss for that specific patient.

**Pre-operative erythrocyte stock.** The pre-operative Hb level is the best predictor of the need for blood transfusion after various elective orthopaedic procedures. 25-29 For primary THR or TKR the risk for the patient of requiring blood transfusion is virtually absent if the Hb level is 14 g/dl or higher, and it increases as the pre-operative Hb level decreases. Patients with a pre-operative Hb level of < 13 g/dl have a four times greater risk of having a transfusion than those with an Hb level between 13 g/dl and 15 g/dl. 27 With a pre-operative Hb level < 11 g/dl, the odds ratio for receiving transfusion is 13.92 compared with non-anaemic patients. 29 A correlation has also been noted between the need for transfusion and gender 29 and body-weight, 27-29 both of which are correlated with the blood volume, and also with advanced age 28, 30 and rheumatoid arthritis 30 owing to the higher prevalence of anaemia. The erythrocyte stock of an individual is related to his or her blood volume and haematocrit. The blood volume in males amounts to 65 ml/kg to 75 ml/kg of body-weight and this is lower in females at 55 ml/kg to 65 ml/kg of body-weight. The difference is related to the differing proportions of less well-perfused adipose tissue. Similarly, an obese male or female will have a smaller blood volume and erythrocyte stock than a lean individual of similar gender and weight.

Various formulae have been proposed to calculate with more precision the blood volume of individual patients, based on their height and weight. 31, 32 In a clinical setting, a simple calculation will give sufficient accuracy for clinical guidance. The erythrocyte volume is calculated from the estimated blood volume and the measured haematocrit. However, as already mentioned, the Hb level is a simple and convenient surrogate for the erythrocyte volume in evaluating the need for transfusion.

**Anticipated blood loss.** The total blood loss associated with an operation can be calculated from the decrease in the haematocrit and Hb level from the pre-operative values to those observed five to seven days after operation, taking into account any transfused blood, as well as the estimated blood volume of that specific patient. Blood loss includes several components. The external blood loss can be measured with reasonable accuracy and includes intra-operative and post-operative measurements, if the wound has been drained. The external blood loss is invariably smaller than the overall calculated blood loss. There is an internal blood loss which is referred to as the ‘hidden’ or ‘occult’ blood loss, which can also be calculated. 33

It is now customary to report the total blood loss in clinical studies, while in the past only the external blood loss was described although there was often sufficient data to allow the total and hidden blood loss to be calculated. Based on such data, it appears that, if no specific blood-saving measures are taken, the external blood loss after THR ranges between 830 ml and 1460 ml, with a calculated total blood loss of between 1550 ml and 2400 ml. 34 The hidden blood loss thus ranges between 720 ml and 840 ml. 34 After total TKR the external blood loss ranges between 570 ml and 1360 ml, and the calculated total blood loss between 1470 ml and 2500 ml, corresponding to a hidden blood loss of 900 ml to 1140 ml. 34 The mean hidden blood loss represents 29% of the total blood loss after THR and 45% to 60% after TKR. 33, 34 The mean hidden blood loss after the surgical treatment of hip fractures was found to range between 547 ml (screw fixation) and 1473 ml (intramedullary nailing). 35

The hidden blood loss has sometimes been attributed to haemolysis, but there is no evidence for this, and scintigraphy using labelled erythrocytes has shown that it is essentially related to the extravasation of blood into the soft tissues. 36

The total blood loss may be higher after revision THR or spinal fusion (up to 2450 ml) or scoliosis surgery (> 3300 ml). 37 Blood loss is procedure-specific, and may also be surgeon- or institution-specific.

In everyday clinical practice, the estimated loss of blood and erythrocyte volume can be no more than a rough estimate, for a number of reasons. First, the patient’s blood volume is estimated rather than measured, and the approximation therefore also affects the estimation of the pre-operative erythrocyte volume. Blood lost intra-operatively has, in theory, a haematocrit similar to the pre-
operative value, although it may become increasingly lower if the patient is receiving large amounts of intravenous fluids in order to maintain normovolaemia. Blood drained post-operatively has a lower haematocrit, which also decreases since it contains an increasing proportion of exudate from the wound, as well as an increasing proportion of free Hb from haemolysis of the extravasated blood. This will nevertheless be listed as ‘blood loss’ although it should more correctly be characterised as ‘bloody fluid loss’. If part of the blood drained post-operatively is re-infused after filtration, this will introduce a source of error in the calculation of total blood loss. Accuracy of the calculations can be improved to some extent by using the mean post-operative haematocrit of the patient. Secondly, there is often post-operative fluid retention for several days, which will lower the post-operative haematocrit while not reflecting the true loss of erythrocyte volume. Finally, if the patient has received a transfusion of red blood cell concentrates a source of error is introduced in the calculation of total blood loss since these ‘red cell units’ are usually considered to provide 200 ml of red blood cells, although their volume may range between 250 ml to 300 ml with a level of haematocrit of between 55% and 80%.

Evaluation of the anticipated blood loss should also take into account pre-existing anomalies of haemostasis, the intake of anti-inflammatory drugs and any factor which influences bleeding.

Acceptable blood loss. The amount of blood loss which a patient can tolerate depends to a large extent on his or her specific physiological features. An elderly patient with coronary heart disease will not tolerate blood loss as well as a younger healthy individual. Therefore, the minimum level of Hb and haematocrit which must be maintained post-operatively should be determined for each patient. For any given level selected, the permissible blood loss will thus depend on the estimated blood volume and pre-operative Hb level and haematocrit. If the anticipated blood loss is higher than the permissible blood loss, some action is necessary.

Methods of reducing blood transfusion

The need for alloegenic blood transfusion may be reduced by various measures including the optimisation of the pre-operative erythrocyte stock, the reduction of blood loss, the lowering of the transfusion trigger, the optimisation of the use of the patient’s own blood and the use of ‘blood substitutes’.

Optimisation of the pre-operative erythrocyte stock. The pre-operative erythrocyte stock seems to be the best predictor of the need for alloegenic transfusion. It should therefore be evaluated sufficiently in advance to allow it to be augmented if necessary. It is estimated that approximately 20% to 30% of the patients who are to undergo THR or TKR are moderately anaemic and are at risk for transfusion if no specific action is taken. Anaemia may be related to iron deficiency, reflected by a low level of plasma iron and elevated transferrin. In such cases, oral administration of iron is commonly prescribed, although it cannot be effective in the short term and is either inefficient, particularly in patients with inflammatory disease, or poorly tolerated in a number of cases. Intravenous administration of iron-sucrose has been used effectively in such cases and is gaining in favour, even in anaemic patients with no demonstrated iron deficiency. In anaemic patients who are not iron- or vitamin B12-deficient, and in the absence of any possible source of chronic blood loss, the Hb level and haematocrit will only improve with the administration of erythropoietin, combined with oral or intravenous administration of iron. Different regimens have been tried in an attempt to achieve optimal cost-efficiency, and two have been advocated, namely, the daily administration of 300 U/kg body-weight for 15 days beginning ten days before surgery or the weekly administration of 600 mg/kg body-weight starting three weeks before surgery with one more injection on the day of surgery. Whichever is used, the erythropoietin must be combined with oral or intravenous iron administration.

Reduction of peri-operative blood loss. This is possible by the attention to a number of details which are now described.

Any congenital or acquired predisposition to bleeding should be detected well ahead of surgery. This includes a variety of congenital conditions, some of which may remain undiagnosed until adult age, such as von Willebrand disease.

The intake of non-steroidal anti-inflammatory drugs (NSAIDs) or aspirin has been associated with increased blood loss. These drugs should be discontinued at least one week before surgery unless this would carry a serious risk in a patient with cardiovascular problems.

Chemical thromboprophylaxis may also influence bleeding. Several meta-analyses have concluded that the risk of major haemorrhagic accidents is similarly low for low-molecular-weight heparins (LMWH), vitamin K antagonists and aspirin, whereas unfractionated heparin gives a higher rate of major haemorrhagic problems. However, bleeding appears to be enhanced even with uneventful thromboprophylaxis using LMWH.

The position of the patient may affect intra-operative bleeding. This is well known in spinal surgery. Bleeding during THR has been reported to be less with the patient in the lateral position than in the supine position, although a recent study has not confirmed this.

Elevation of the lower limb after TKR has been shown to reduce post-operative blood loss.

Controlled hypotensive spinal or epidural anaesthesia has been shown to reduce blood loss compared with general anaesthesia. Maintaining normothermia is also essential to avoid excessive bleeding. Pre-operative arterial embolisation is reserved for specific indications, such as complex pelvic fractures or surgery on highly vascularised tumours such as metastases from renal-cell.
cancer or aneurysmal bone cysts. Careful surgical haemostasis is recommended, with the occasional use of bone wax and surgicel when indicated.

Several studies have shown that post-operative bleeding can be reduced by half by using local haemostatic agents such as fibrin sprays. They have demonstrated their efficacy in a variety of surgical operations for more than 30 years. The limiting factor is essentially economic, owing to their high cost, although a low-cost locally-prepared fibrin glue has also been found to be highly efficient. Local application of plateleter gel before wound closure is another option, and reduces bleeding after TKR. The topical use of tranexamic acid has also been shown to reduce blood loss after various operations such as coronary bypass or screw fixation of the lumbar spine.

Systemic haemostatic agents have been used in an attempt to decrease blood loss and transfusion requirement. Desmopressin, a synthetic vasopressin-analogue, has been used with variable success in cardiac surgery. Several clinical trials failed to show any difference in blood loss and transfusion requirements when desmopressin was used in elective THR or TKR in haematologically normal patients. A Cochrane systematic review concluded that there was no convincing evidence that desmopressin reduced allogenic blood transfusion in patients who did not have congenital bleeding disorders. High-dose aprotinin, a fibrinolytic inhibitor, was found to reduce blood loss in primary and revision THR, TKR and spinal fusion. Its use has been limited because of its high cost and possible allergic reactions and a recently reported risk of acute renal failure. The worldwide sale of Trasylol (Bayer, Leverkusen, Germany) was suspended in 2007 pending the results of a trial to evaluate its benefit-risk ratio. Tranexamic acid, a synthetic fibrinolytic inhibitor, has been used for more than 20 years in various fields such as dentistry, gynaecology, cardia surgery, urological surgery, and liver transplantation. Several studies have shown that it reduces total blood loss by nearly 50% and markedly reduces the need for allogenic blood transfusion after THR, TKR and spinal surgery. It is relatively inexpensive and does not appear to have any significant untoward side-effects. It did not increase the incidence of venous thrombosis in any of the clinical studies reported. For greatest efficacy it should be administered at the beginning of surgery, either as a continuous perfusion or as repeated bolus injections until a few hours after operation, because its half-life in the plasma is short. A comparative prospective, randomised controlled study showed a similar reduction in blood loss with the local application of fibrin spray and the intravenous administration of tranexamic acid. Similar beneficial effects have been reported with the oral administration of tranexamic acid before and up to 18 hours after TKR. Recombinant Factor VII has been found to be beneficial in controlling coagulopathic bleeding in massively bleeding surgical or trauma patients, but it has no indication in elective orthopaedic surgery since its cost is too prohibitive.

Many studies have investigated the possibility of reducing blood loss by not using wound drainage. Although wound drainage is an established tradition in orthopaedic surgery, there is considerable evidence to support its discontinuation in a number of common operations. In a recently updated meta-analysis, Parker et al concluded that there was insufficient evidence to support the routine use of closed suction drainage in orthopaedic surgery. Pooling data from randomised or quasi-randomised comparative studies, they failed to note any significant difference in the incidence of wound infection, haematoma, dehiscence or re-operations between drained and undrained wounds. Blood transfusion was required more often in patients who had drains, while bruising was more common in those without drains, who also needed more frequent dressing reinforcement. Differing results have been reported with respect to the effect of non-drainage on blood loss and transfusion rates, which is not surprising considering the number of variables involved, such as the number and diameter of drains, the type of suction used, the time to removal of the drain, and the concomitant use of LMWH or NSAIDs. The use of transfusion rates and volumes as outcome measures may also be biased by local differences in transfusion practice. Some authors have reported a reduction of blood loss and transfusion requirements after non-drainage, while others reported no difference. The avoidance of drainage reduces the external blood loss, but not necessarily the hidden loss, which may be increased, particularly after TKR. However, the overall balance appears to be towards a reduction in total blood loss. Practical details can make a real difference, such as an efficient padded compressive dressing which will reduce bleeding through a tamponade effect. This is common practice after operations on the knee, but it has also proved efficient at the hip.

Various modifications in the drainage technique have been tested, such as delayed or intermittent drainage. These have usually resulted in a reduction in external blood loss by up to 50%, but not in hidden loss. In one study on THR no reduction in external blood loss was noted after clamping the drain for the first two hours, while another reported marked reduction in external and total blood loss following intermittent clamping during the first six hours, with the drain opened for five minutes every hour. Among numerous trials on TKR only one reported no reduction in blood loss with clamping of the drain for the first two hours. Other trials showed a significant reduction in external blood loss using delayed drainage with the drain opened after one or four hours or intermittent drainage with the drain opened for five or ten minutes every one or two hours for the first six to ten hours. Delayed or intermittent drainage, as with non-drainage, relies on a tamponade effect. In other studies, delayed drainage was combined with the injection of 30 ml or 50 ml of saline with dilute adrenaline (1:200 000 to 1:1 000 000) into the knee after TKR. This also effectively reduced the external blood loss.
loss. Problems of wound healing were noted only when adrenaline was left in situ for 24 hours. Infiltration of soft tissues with dilute adrenaline before closure is commonly used without similar adverse effects.

The usual justification for systematic wound drainage is the fear of encouraging the formation of deep haematoma if the wound is not drained. However, sonography and scintigraphy with labelled red blood cells have shown that haematomas and seromas will form anyway after removal of the drain. Drains should be removed after 24 hours, since 90% of the blood is collected within the first 24 hours and the limited amount of blood and/or exudate which could be collected thereafter would not balance the rising risk of retrograde infection.

The use of the pneumatic tourniquet increases the comfort of the surgeon and is often instinctively perceived as improving cement fixation in TKR. However, it does not reduce the total blood loss and it may even increase it. A prolonged tourniquet time may induce a post-ischaeamic reperfusion injury resulting in reactive hyperaemia and oedema. It may also stimulate fibrinolysis and increase the hidden blood loss.

The timing of deflation of the tourniquet remains contentious. Deflation before wound closure tends to increase blood loss, while deflation after wound closure risks an untoward. Deflation before wound closure tends to increase oedema. It may also stimulate fibrinolysis and increase perfusion injury resulting in reactive hyperaemia and prolonged tourniquet time may induce a post-ischaemic reperfusion injury.

Hypervolaemic haemodilution does not involve the use of medicinal drugs and of haemostasis if extreme haemodilution is possible using a variety of techniques. The use of the pneumatic tourniquet increases the comfort of the surgeon and is often instinctively perceived as improving cement fixation in TKR. However, it does not reduce the total blood loss and it may even increase it. A prolonged tourniquet time may induce a post-ischaeamic reperfusion injury resulting in reactive hyperaemia and oedema. It may also stimulate fibrinolysis and increase the hidden blood loss.

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Lowering the transfusion trigger. As already mentioned, the historical transfusion trigger of 10 g/dl of Hb and 30% haematocrit can safely be lowered in most patients undergoing elective orthopaedic operations. Currently, accepted thresholds are 8 g/dl of Hb in the older patient without severe comorbidities, and 7 g/dl in the younger healthy patients. This should of course be tailored to the individual patient.

Optimisation of the use of the patient’s own blood. This is possible using a variety of techniques.

Acute isovolaemic haemodilution presents, in theory, a number of advantages. Shed blood has a low haematocrit, which reduces red blood cell loss. Haemodilution enhances microcirculation. Autologous blood with active platelets and coagulation factors is re-infused and the risk of admin-istrative error is virtually non-existent. However, it is a time-consuming procedure. Some patients tolerate acute blood withdrawal poorly and it also carries its own risks such as ischaemia if the target haematocrit is very low. In addition, there are the problems of re-transfusion of anaesthetic drugs and of haemostasis if extreme haemodilution is achieved. Hypervolaemic haemodilution does not involve any withdrawal of blood and is less time-consuming. It has been found to have similar efficacy in reducing the need for allogenic blood transfusion in healthy adults, but it shares most of the disadvantages of isovolaemic haemodilution and can only be considered in young healthy individuals with efficient physiological compensation mechanisms.

A meta-analysis of 24 randomised trials concluded that the impressive reduction in allogenic blood transfusion reported in many instances may in fact be related to flawed study design. In a comparative study of the blood-sparing effect of tranexamic acid versus isovolaemic haemodilution, Zohar et al concluded that both were effective but that tranexamic acid appeared to be preferable because it was associated with superior haemodynamic stability and was also better in terms of blood sparing.

Pre-operative autologous blood predonation also offers a number of theoretical advantages. It has been widely used in the recent past, but has now fallen into relative disfavour for several reasons. It cannot reasonably be considered in patients who are anaemic, i.e. in 20% to 30% of patients who are to undergo THR or TKR, or in those with an Hb level > 14.5 g/dl since they will not require blood transfusion. It should therefore be targeted to men with an Hb level of 11.0 g/dl to 14.0 g/dl and to women with a level of 13.0 g/dl to 14.0 g/dl whose anticipated peri-operative blood loss is close to 1000 ml. In older patients who are not anaemic, the capacity of the erythropoietic marrow to react to the stimulus of blood withdrawal may prove to be inadequate. They may come to operation with an Hb level lower than that before pre-donation, and will need more transfusion episodes (autologous or allogenic) than they would have otherwise needed. For the same reason, many patients simply cannot complete the pre-donation programme because they become increasingly anaemic. The incidents and complications related to blood pre-donation are nearly 12 times as likely in autologous blood donors than in blood-bank donors, and they may be associated with significant morbidity.

Pre-operative autologous blood predonation has been shown to reduce the absolute risk for allogenic blood transfusion by 43.8%, but at the same time to increase the risk of any transfusion (allogetic or autologous) (relative risk: 1.29). The proportion of pre-donated blood which is wasted is high, up to 50% in some studies. However, Mercuriali and Inghiller achieved a wastage rate of < 15% with customised prediction of transfusion requirements for the individual patient. It is generally accepted that pre-donated blood should not be re-infused in the absence of a clear indication and should also not be used for transfusion in other patients.

Brecher and Goodnough therefore concluded that “judicious use of pre-operative donation in concert with other blood conservation methods in specific cases remains appropriate, however automatic referral of all patients for pre-operative donation is oversimplistic and should be discouraged”. An increasing number of studies comparing pre-operative donation and pre-operative administration of erythropoietin have concluded that the latter appears to be better with respect to the prevention of post-operative anaemia, while its cost-efficiency also appears to be superior when direct and indirect cost of pre-operative donation are taken into account. Pre-operative donation can
also be combined with the pre-operative erythropoietin. This enables some patients to complete a pre-donation programme which would otherwise not have been possible, but this combination appears to be useful only when the anticipated blood loss is particularly heavy.

Another blood-saving procedure is peri-operative blood salvage. This includes re-infusion of blood drained within six hours after operation using cell salvage. Blood shed during operation may be altered by irrigation fluid, air and cement and should therefore be washed before re-infusion to avoid coagulopathy.

Peri-operative blood salvage has been shown to be beneficial in cases of blood loss exceeding 1000 ml, but is not considered to be cost-effective in primary arthroplasty if other effective blood-saving measures have already been taken. Its use is generally restricted to specific indications in which the anticipated blood loss is very high, such as revision THR. Carless et al noted in their Cochrane Systematic Review that the use of cell salvage reduced the rate of allogenic transfusion by a relative 39% (relative risk 0.61), with a mean saving of 0.67 units of allogenic blood per patient. However, at least two units need to be recovered for the method to be cost-effective. In an extensive study on cost-effectiveness of various blood saving techniques, Davies et al concluded that cell salvage (intra- and/or post-operative) had lower costs and gave slightly higher quality-adjusted life years than all of the alternative transfusion strategies except acute normovolaemic haemodilution. However, their analysis was not directed specifically at orthopaedic surgery. They suggested that post-operative re-infusion of unwashed cells may be more cost-effective than intra-operative salvage in orthopaedic procedures.

The available salvage and re-infusion systems use either washed or unwashed filtered cells. Most re-infuse unwashed cells after filtration and the specific features of the filter used in each system may result in the re-infusion of a variable amount of leukocytes and red blood cells. Blood re-infused after simple filtration has a low haematocrit and contains large concentrations of inflammatory mediators, histamine, eosinophil cationic protein, eosinophil protein X, myeloperoxidase, plasminogen activator inhibitor type 1, activated complement factor C3 and various coagulation factors and split products, as well as free Hb. Despite this, post-operative re-infusion of unwashed salvaged blood from drains appears to be safe, with no clinically important side-effects when a low volume of salvaged blood is returned. It is generally accepted that no more than 1000 ml of drained blood should be re-infused, and also that blood drained more than six hours after the end of surgery should not be re-infused.

Post-operative cell salvage is not always cost-effective since the volume of blood recuperated is unpredictable. The reported mean volume of re-infused blood has ranged between 360 ml and 880 ml, relating to differences in surgical or drainage technique (drain number and diameter, vacuum pressure, etc); there are also marked variations among patients. The method is useful in patients with a small blood volume, as the likelihood of having to re-infuse more than 10% of their blood volume is high. It can be used in combination with other blood-saving measures if the anticipated blood loss is high, or in patients with a small blood volume and a low pre-operative Hb level. Most but not all studies on post-operative cell salvage have reported a reduction in allogenic transfusion rate and volume.

Intra- and post-operative cell salvage are contraindicated in patients with infection and malignancy. The use of ‘blood substitutes’. Since World War II, there has been a search for possible blood substitutes which would be readily available in emergency situations. The perfect ‘blood substitute’ should have an oxygen-carrying capacity similar to that of Hb, should not require cross-matching, have a long shelf life as well as a long intravascular half-life, and be free from side effects. Two main types of possible blood substitute still in development have reached the stage of clinical investigation, namely perfluorocarbons and haemoglobin-based oxygen carriers.

Perfluorocarbons are biologically inert volatile fluids with a high dissolving capacity for O2 and CO2. They can transport and deliver oxygen by simple physical dissolution. Since they are not water-soluble, they are administered as emulsions stabilised by phospholipids, which can be sterilised. Their O2 carrying capacity depends on the concentration of perfluorocarbons, which varies between products.

Haemoglobin-based oxygen carriers are based on natural or recombinant human Hb or bovine Hb, and are produced by various procedures including purification, encapsulation in synthetic phospholipid liposomes, cross-linkage and polymerisation in order to limit the possible side-effects of free Hb. The clinical development of haemoglobin-based oxygen carriers has indeed been slowed down by adverse effects, including an increase in blood pressure by systemic vasoconstriction due to the scavenging of vascular endothelial nitric oxide, and oxidation, which generates methaemoglobin. Haemoglobin-based oxygen carriers have been tried in pre-clinical and clinical studies but no product has yet achieved market approval in Europe or the United States. One of the limitations of Hb solutions is their short intra-vascular half-life, and this is why they are intended to serve as a bridge to transfusion in emergency situations, rather than as definitive blood substitutes. There is no indication that they could be used in elective orthopaedic operations, at least in the near future.

Conclusions

Blood management is a complex issue, and most surgeons cannot be expected to have the necessary time and expertise to select from the numerous blood-saving techniques available to determine the optimal combination for each individual patient. Blood management should be made as simple as possible and should rely on common sense while also being cost-effective and having a low risk of complica-
tions. Choices are made more complicated by the contradictory results reported with most blood-saving techniques. Critical analysis of the vast literature on blood management clearly shows the poor methodological quality of most studies. There are few prospective, randomised controlled studies, and frequent methodological flaws related to inadequate blinding and to the use of subjective outcome measures such as the rate and amount of allogeneic blood transfused. With respect to transfusion policies there are major variations between institutions as there are with anaesthetics, surgical and wound-drainage techniques. Allogeneic transfusion rates reported in the literature in the absence of specific blood-saving measures after TKR have ranged from 6% to 95%. In a study comparing blood loss after TKR with and without computer-assisted techniques, a mean blood loss of 1700 ml in the drains was reported in the control group, a figure which would be found alarming by a number of surgeons with an interest in blood management, and the authors’ conclusion that computer-assisted TKR reduces blood loss may not apply to patients managed using a different blood management policy. Similar scepticism also applies to the reported blood-saving effect of mini-invasive hip or knee replacement, for which the evidence appears to be very small.

All meta-analyses of studies on blood management, including Cochrane Systematic Reviews, have reported that the level of evidence is low for most conclusions and statements in the literature. They invariably conclude that there is a need for large prospective, randomised controlled studies, but these may in fact never be done at all. Common sense therefore should prevail. Several easy safe and inexpensive measures can definitely help to reduce the rate and volume of allogeneic blood transfusion. First, patients should be evaluated at least one month before surgery, to check their Hb and haematocrit levels, and to make sure that they do not have an increased risk of bleeding. The surgeon should then consider the anticipated blood loss, should be aware of the existence of a hidden blood loss, and estimate whether this is not higher than the allowable blood loss in each patient, which requires an evaluation of the erythrocyte stock. If the anticipated blood loss is higher, some action is necessary. Pre-operative blood donation has been used as a sort of panacea, but has often been found to be counterproductive since it may make the patient more anaemic and increase the risk for transfusion. Erythropoietin and iron administration now appear to be the best choice for patients with a Hb level <11.0 g/dl and an anticipated blood loss of 1000 ml to 1500 ml such as in primary THR or TKR. With a pre-operative Hb level at 12 g/dl or 13 g/dl, simple measures such as the systemic administration of tranexamic acid, local application of fibrin spray, non-drainage of the wound or delayed and intermittent drainage, or post-operative cell salvage may be sufficient to avoid allogenic blood transfusion. These measures are not mutually exclusive and have all been found to be effective, although with the usual low level of evidence already mentioned. In cases of high anticipated blood loss, pre-operative blood donations may be considered, possibly combined with the administration of erythropoietin, in addition to all the other measures and with the addition of intra-operative cell salvage.

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