Platelet transfusion

The statement printed below was agreed at a consensus conference on platelet transfusion organised by the Royal College of Physicians of Edinburgh and held in Edinburgh in November 1997. We publish this statement at the request of the organising committee to bring it to the attention of physicians who do not read the haematological literature. The statement will also appear in the British Journal of Haematology in 1998 with the scientific evidence upon which it is based.

Platelet transfusion is used in patients with a low platelet count (thrombocytopenia) or disordered platelet function who are actively bleeding (therapeutic use) or who are at serious risk of bleeding (prophylactic use). There is extensive clinical evidence that platelet transfusions are valuable but the procedure also carries risks and costs. The use of platelet transfusion has developed without much clinical trial-based evidence on its effects and complications, or on the value of additional procedures such as leucocyte depletion. This has obvious implications for the ethically crucial questions of patient safety and the balance of risks and costs to benefits. Despite the difficulties in performing randomised clinical trials on an established procedure, the conference is firmly of the view that such studies are needed if clearer, evidence-based guidelines for the use of platelet transfusion are to be formulated. Even where randomised trials are not possible, the conference recommends that all decisions to transfuse platelets should be made according to written institutional protocols that will allow generally applicable conclusions to be drawn from the results.

Indications for platelet transfusion

Prophylactic

The main use of platelet transfusion is in the prevention of bleeding in patients with haematological malignancies (particularly leukaemias) who have bone marrow failure caused by their disease or its treatment.

For patients with bone marrow failure, it has been accepted practice to transfuse platelets where levels are very low. Based on the evidence presented at the meeting there was a general agreement that a platelet threshold of 10 x 10^9/L is as safe as higher levels for treating most patients without additional risk factors. These risk factors – which include sepsis, concurrent use of drugs (eg antibiotics) and other abnormalities of haemostasis – are indications for a higher threshold. Higher threshold numbers are also needed to cover invasive procedures, eg line insertions and biopsies, but there is no consensus on appropriate thresholds. For uncomplicated patients, evidence on the safety of even lower platelet thresholds than 10 x 10^9/L should be sought. However, accurate counting of low platelet numbers may create difficulties when trying to reduce the threshold below 10 x 10^9/L. In neonates where there is a considerable danger of haemorrhage, platelet transfusion is indicated at a higher threshold than in adults.

The avoidance of low haematocrit in patients with thrombocytopenia or disordered platelet function reduces the risk of haemorrhage.

Therapeutic

Most major surgery (including cardiac and vascular) can be successfully carried out without platelet transfusions. Patients who have taken aspirin in the ten days prior to surgery have increased risk of bleeding and the value of continuing aspirin at this time should be evaluated.

In massive haemorrhage, the first priorities are to achieve surgical haemostasis and resuscitation. There is a consensus to transfuse platelets if the count is less than 50 x 10^9/L, but it is clear that other clinical criteria need to be considered.

Liver transplantation gives special problems with haemostasis, and platelet transfusion is frequently required. In liver surgery, thromboelastography has been shown to be a good predictor of platelet need and deserves evaluation in other areas. Antifibrinolytic agents have a platelet-sparing role in liver transplantation and complex cardiac surgical operations. They may also be used in other situations, as may agents such as DDAVP, when platelet numbers or function are compromised.

Platelet transfusion is also used to treat clinically significant bleeding in patients with idiopathic thrombocytopenic purpura (ITP). The consensus view is that platelet transfusions have been over-used in this area, especially in children. Intracranial or eye haemorrhage or severe bleeding from the gut are the major concerns; except in these circumstances platelet transfusion should be avoided.

Risks and contraindications

The most common acute risk is bacterial infection which has been under-reported. The important long-term risks of transmitting viral infection from the donor are well recognised and have been substantially reduced. However, the threat from newly recognised viruses is always present. The theoretical possibility that blood products may be able to transmit the agent of variant CJD has attracted attention and cannot be discounted. Experimental procedures for reducing the risk of transmitting bacterial and viral infection by chemical treatment of the platelet preparations (eg by psoralens plus ultra-violet light) are under investigation.

The presence of white blood cells in the platelet preparation enhances the risk of certain viral infections and is largely responsible for generating cytokines that can give
rise to febrile reactions after platelet transfusion. Platelet transfusion through negatively charged filters can also activate the contact system and give rise to significant hypotension in patients receiving ACE inhibitors. A ten-fold reduction in white cell numbers (leuco-reduction) before storage of platelets is sufficient to abolish most febrile reactions, and is readily achievable in both buffy coat derived and apheresis platelets. White cell numbers can be further reduced to less than 5 x 10^6 per dose (leuco-depletion) by specialised manufacture, by improving the apheresis technique and by pre-storage filtration. Leuco-depleted platelets reduce transmission of some viruses and febrile reactions, as well as reducing HLA allo-immunisation which can cause patients to become refractory (resistant) to further platelet transfusions.

Irradiation of platelet concentrates is required to prevent graft versus host disease and this needs to follow national guidelines.

Platelet transfusion is contra-indicated in heparin-induced thrombocytopenia, thrombotic thrombocytopenic purpura and the haemolytic uraemic syndrome.

**Recipients who are refractory to platelet transfusions**

Failure to achieve a satisfactory response to platelet transfusion (refractoriness) occurs in up to half of those receiving prophylactic transfusions. This is defined by a poor increment in platelet count rather than on clinical grounds. Patients who remain refractory for immunological reasons often receive prophylactic transfusions with HLA-matched or crossmatch-compatible platelets. Satisfactory increments are frequently obtained by these approaches but their effectiveness in reducing severe bleeding deserves more detailed evaluation – as do the relative effectiveness of HLA-matching compared to platelet crossmatching, and the issue of whether patients who remain refractory despite these measures should only be transfused therapeutically.

**Quality control of platelet preparations**

In addition to those already routinely undertaken, quality control measures should include: more accurate measurements of low white cell and high platelet counts; more sensitive tests for microbiological safety; and rapid techniques for detecting viral nucleic acid in platelet preparations. There are few appropriate surrogate markers that are relevant to the arrest or prevention of bleeding in patients, although the swirl test has its adherents.

Performance in quality assurance should meet evolving national standards.

**Conclusion**

The overall conclusion is that platelet transfusion is a well established clinical procedure, but that it can never be entirely safe and must be given only where there is clear clinical justification. Nevertheless the precise indications for its use and the optimal specification of the product still need to be defined. These goals will be achieved only by a combination of prospective randomised trials with the information that will come from the use of clearly defined protocols, and their efficient audit.

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**Members of the consensus panel:**

Professor Alan K Burnett (Vice-Chairman), Dr Virginia Clough, Ms Maggie Grundy, Mr Chris Holme, Professor Peter J Lachmann (Chairman), Mr Charles Marshall, Mr Lloyd Scott, Dr Patricia Skacel, Dr Colin J Sinclair, Mr Peter C Taylor, Mrs Pat Walsh.

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**Inflammation, genes and gene therapy**

The Joint Symposium of the three Royal Colleges of Physicians was held on 20 November 1997 in Aberdeen to mark the opening of the Institute of Medical Sciences at the University of Aberdeen.

**Inflammatory disease**

Anti-inflammatory treatment for asthma

Professor Peter Barnes (Imperial College School of Medicine at the National Heart and Lung Institute) described the inflammatory basis of asthma and the best way to treat the disorder. Even mild asthma is a chronic inflammatory condition marked by infiltration of the airway with activated eosinophils, T cells, macrophages and degranulating mast cells that leads to epithelial shedding. The epithelial cell is a key mediator in this inflammatory response, producing a range of inflammatory mediators including pro-inflammatory cytokines (interleukin-1 (IL-1), tumour necrosis factor (TNF)), chemokines (IL-8, eotaxin) and vasodilators (nitric oxide, prostaglandin E2). This cell is also the prime target of inhaled steroids, the mainstay of anti-inflammatory treatment in asthma. Steroids bind to the glucocorticoid receptor in the cell, which then binds to and inhibits transcription factors such as NF-κB that are pivotal in activating synthesis of pro-inflammatory cytokines and enzymes. Inhibiting the inflammatory activity of the epithelial cell affects the injury throughout the thickness of the airway wall.

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