Guidelines for the diagnosis, investigation and management of polycythaemia/erythrocytosis

Mary F. McMullin,1 D. Bareford,2 P. Campbell,3 A. R. Green,3 Claire Harrison,4 Beverley Hunt,4 D. Oscier,5 M. I. Polkey,6 J. T. Reilly,7 E. Rosenthal,8 Kate Ryan,9 T. C. Pearson4 and Bridget Wilkins10 On behalf of the General Haematology Task Force of the British Committee for Standards in Haematology

1Department of Haematology, Queen's University, Belfast, Belfast City Hospital, Belfast, 2Department of Haematology, City Hospital, Birmingham, 3Department of Haematology, University of Cambridge, Cambridge Institute for Medical Research, Cambridge, 4Department of Haematology, St Thomas' Hospital, London, 5Department of Haematology, Royal Bournemouth Hospital, Bournemouth, 6Sleep and Ventilation Service, Department of Respiratory Medicine, Royal Brompton Hospital, London, 7Department of Haematology, Royal Hallamshire Hospital, Sheffield, 8Consultant Paediatric Cardiologist, Guy's Hospital, St Thomas Street, London, 9Department of Clinical Haematology, Manchester Royal Infirmary, Manchester, and 10Cellular Pathology Department, Royal Victoria Infirmary, Newcastle upon Tyne, UK

Keywords: erythrocytosis, polycythaemia, polycythaemia vera, secondary erythrocytosis, management.

Traditionally, polycythaemia has been used to identify a group of varied disorders with an increase in circulating red cells that are typified by a persistently raised haematocrit (Hct). Since only the red cell lineage is involved, the term erythrocytosis has more validity and will be used throughout this article. Polycythaemia will be retained in relation to the clonal disorder, polycythaemia vera (PV), in which three cell lineages are involved.

Aim

The purpose of this guideline is to provide a rational approach to the diagnosis, investigation and management of patients with an erythrocytosis. This will include recommendations on the management of PV, apparent and relative erythrocytosis, idiopathic erythrocytosis and the secondary erythrocytoses because of high oxygen affinity haemoglobin, hypoxia because of chronic lung disease, congenital cyanotic heart disease and postrenal transplantation.

Methods

The guideline group was selected to include UK-based medical experts. The drafting group met (real or virtual) on four occasions and communicated by e-mail. Each member of the group was allocated responsibility for the preparation of a selected component of the first draft. Medline, CANCERLIT and EMBASE were systematically searched for publications in English from 1966 to June 2004. Relevant literature in group members own collections and older references generated from initial papers were also examined. The Cochrane controlled trials register and the Cochrane optimal search strategy for randomised controlled trials was searched but no additional material was identified. Randomised trials and series of patients and single case reports were considered if appropriate. Meeting abstracts were not included in the systematic search strategy. The group leader synthesised the draft components, which were subsequently revised by consensus. No recommendations are included for which full consensus was not achieved. The guideline was reviewed by sounding boards, and British Committee for Standards in Haematology (BSCH) and comments were incorporated where appropriate. Criteria used to quote levels and grades of evidence are outlined in Table I.

Diagnosis of erythrocytosis

Patients with a persistently raised venous haematocrit (Hct) (>0.52 males, >0.48 females for >2 months) should, in general, be investigated by measurement of their red cell mass (RCM). A number of physiological factors have been shown to influence the Hct value, although in practice the use of only minimal or no venous occlusion when taking the blood sample is the most important. In addition, Coulter ‘S’ and Coulter ‘S Plus’ analysers are known to underestimate Hct [approximately 7% at reduced mean cell haemoglobin (MCH) values]. As a result, a correction factor should be applied (Guthrie & Pearson, 1982).

However, males and females with Hct values above 0.60 and 0.56 respectively can be assumed to have an absolute erythrocytosis and do not require confirmatory studies.
The investigation of absolute erythrocytosis

The classification of absolute erythrocytoses is shown in Table II. Once an absolute erythrocytosis has been confirmed it is desirable to identify the underlying aetiology, although this may not be possible either initially or after prolonged investigation. Nevertheless, the starting point is knowledge of the underlying causes of a secondary erythrocytosis (Table II) and the diagnostic criteria of PV (Table III). Frequently, it can be difficult to prove conclusively that an erythrocytosis is either secondary or primary and dual pathologies resulting in an erythrocytosis should be considered, especially in the elderly. Two stages of diagnostic tests are listed in Table IV (see below for comment). The number and order of tests and intensity of investigation depends on the clinical features after assessment. However, stage 1 investigations, including serum erythropoietin (EPO) levels and blood gas measurements, should be undertaken in all patients and can facilitate the diagnosis in the majority of cases. Selected tests in stage 2 are undertaken following an evaluation of the initial results.

Stage 1 investigations

Full blood count

A neutrophilia is present in approximately two-thirds and thrombocytosis in 50% of PV cases (Berlin, 1975) and, as a result, provide useful minor criteria for the diagnosis of PV (Pearson & Messinezy, 1996). It should be noted that smokers have significantly higher neutrophil counts than non-smokers.
Whitehead et al., 1995) and it has been suggested that, in smokers, the upper limit of normal for neutrophil count should be taken as $12.5 \times 10^9/l$. Serum ferritin and vitamin B$_{12}$

Low serum ferritin levels are more commonly seen in PV than secondary erythrocytosis. Indeed, the absence of iron stores is a frequent finding in PV marrow histology. Although an elevated B$_{12}$ level is characteristic of PV, resulting from transcobalamin release from an increased granulocytic mass, it is not an essential investigation.

Renal and liver function

Erythrocytosis may be associated with both renal and hepatic disease. Serum calcium levels should be determined to exclude the very rare secondary erythrocytosis caused by parathyroid adenomas or carcinoma.

Arterial oxygen saturation

The measurement of arterial oxygen saturation (SaO$_2$), a sensitive indicator of tissue hypoxia, is most easily achieved with the use of a pulse oximeter. However, there are three situations causing hypoxic erythrocytosis in which the SaO$_2$
can be misleading: carbon monoxide poisoning, the presence of high oxygen affinity haemoglobins and sleep apnoea syndrome. Most instruments provide carbon monoxym haemoglobin (COHb) measurements and this value should be subtracted to give an accurate SaO2 result. Smokers generally have higher COHb levels, although a secondary erythrocytosis because of smoking alone is uncommon (Smith & Landaw, 1978). High oxygen affinity haemoglobins, as well as congenitally low 2,3-bisphosphoglycerate levels, will give rise to a normal SaO2, despite tissue hypoxia and measurement of the p50 is important to exclude these rare conditions. A SaO2 below 92% has been taken to indicate a causal relationship with an absolute erythrocytosis (Berlin, 1975). A normal daytime SaO2 can also be seen in the sleep apnoea syndrome and supine hypoventilation because of premature airway closure. It is important therefore to consider these conditions and to enquire about symptoms relating to nocturnal oxygen desaturation, for example snoring, nocturnal restlessness and daytime somnolence. Nocturnal reduction in SaO2, usually but by no means exclusively seen in obesity (Pearson & Treacher, 1990), may be found in 10–20% of patients who would otherwise have been classified as having idiopathic erythrocytosis (Moore-Gillon et al, 1986). Review by a chest or sleep physician (to assess the need for a respiratory sleep study) is recommended in patients with erythrocytosis of all types who are known to snore heavily and either have unwanted daytime somnolence [defined as an Epworth Score >10/24 (www.stanford.edu/~dement/epworth.html)] (Johns, 1993) or who are significantly overweight (body mass index >30 kg/m²). A chest X-ray is also recommended to exclude lung pathology.

**Serum erythropoietin levels**

Since erythrocyte production is controlled by EPO, a serum EPO level can provide information as to whether the erythrocytosis is hormonally mediated or autonomous. In patients with erythrocytosis secondary to hypoxia, serum EPO levels are typically raised. In contrast, EPO levels in patients with PV are characteristically reduced and remain low in the majority of cases even following adequate venesection (Messinezy et al, 2002a). EPO values below the reference range, however, can be seen in idiopathic erythrocytosis, a fact that lowers the specificity of low EPO levels for PV. In addition, a normal serum EPO level excludes neither hypoxia nor PV as the cause of erythrocytosis. Nevertheless, with the availability of specific, sensitive and reproducible EPO assays a low serum EPO levels can now be used as a minor criterion in the diagnosis of PV (Messinezy et al, 2002b).

**Abdominal ultrasound**

Abdominal ultrasound is an essential investigation in all patients with a proven absolute erythrocytosis to exclude underlying renal and hepatic pathology. In the absence of liver disease, a palpable spleen is a reliable sign of PV and, as such, has been adopted as a major criterion for its diagnosis. Splenomegaly can be found in two-thirds of PV cases by various imaging techniques, although ultrasound is the simplest (Messinezy et al, 1997). However, in view of the significant inter-observer error of scanning detection of non-palpable splenomegaly, it has been proposed that this finding be taken as a minor criterion (Table III). In females, pelvic ultrasound would detect leiomyomas, which have occasionally been found to be a cause of secondary erythrocytosis.

**Stage 2 investigations**

**Bone marrow examination**

Bone marrow aspirate and trephine biopsy are not required to meet the diagnostic criteria for PV (Table III). However, these investigations provide useful information including confirmation of the diagnosis of PV, differentiation from secondary erythrocytosis and other myeloproliferative disorders (MPDs), and assessment of the degree of fibrosis. They also provide a baseline which can be compared with subsequent bone marrow examinations to assess disease progression or response to therapy. The finding of a chromosomal abnormality establishes clonality.

In PV, aspirated bone marrow is expected to have dense particles and cellular trails. There is usually marked erythroid hyperplasia with moderate to marked hyperplasia of granulopoiesis and megakaryopoiesis. As well as increased cells of the neutrophil lineage, eosinophils and basophils, but not monocytes, are increased. Wide variations in megakaryocyte size, including larger variants with hyperlobated nuclei are characteristic. Iron stores are typically absent.

The bone marrow trephine core is hypercellular for the patient’s age. There is trilineage involvement, not selectively and only rarely preferentially erythroid (Pierre et al, 2001). Erythroid maturation is maintained and is normoblastic, although erythropoietic nests may be abnormally located abutting trabeculae. Granulocyte maturation is also maintained. It may be left-shifted and distributed in a disorderly fashion, involving loss of the usual preferential localisation of promyelocytes and myelocytes at trabecular margins. There is increased variation in megakaryocyte cell size. Large variants are often predominant and may have uneven or reduced nuclear lobulation. Clusters of megakaryocytes are common and typically pleomorphic, i.e. contain mixed large and small cells. There is usually a mild to moderate increase in stromal reticulin (grade 2–3) (Imbert et al, 2001).

Histological features supporting a diagnosis of secondary erythrocytosis rather than PV include the presence of stromal inflammatory features, such as increased plasma cells, increased haemosiderin in stromal macrophages and evidence of abundant background apoptotic activity (Thiele et al, 2001a).

Bone marrow histology is a B criterion in the WHO classification and is helpful in distinguishing PV from reactive
conditions with secondary erythrocytosis. However, there is histological overlap with other chronic myeloproliferative disorders (CMPD), which currently limits the usefulness of histology in sub-classification (Pierre et al., 2001). (In doubtful diagnostic situations, haematologists or pathologists may find it helpful to refer the sample to a haematopathologist with expertise in trephine biopsy interpretation.)

**Karyotype**

Cytogenetic abnormalities are found in 10–20% of patients with PV. The abnormalities trisomy of chromosomes 8 and 9, del (20q), del (13q) and del (1p) are the most commonly found. An abnormal karyotype, a clonality marker, is a major diagnostic criterion. Patients who progress to myelodysplastic syndrome or acute leukaemia almost always have a karyotypic abnormality (Pierre et al., 2001).

**Culture studies of BFU-E (erythroid burst-forming units)**

The culture of the mononuclear non-adherent fraction of peripheral blood cells or bone marrow cells of patients with PV in serum-containing medium without the addition of EPO leads to the growth of BFU-E – so-called ‘endogenous erythroid colonies’. This feature, which is not usually found in normal individuals or patients with secondary erythrocytoses, has been shown to be a good marker for PV (Partanen et al., 1989). Despite these observations, culture techniques are not standardised and are expensive. It is for these reasons that the finding of endogenous erythroid colonies is best regarded as a minor diagnostic criteria.

**Oxygen dissociation curve (p50)**

It is important to examine the p50 in patients with unexplained erythrocytosis to exclude a high affinity haemoglobin. There are a large number of beta-chain haemoglobin variants that have increased oxygen affinity and a resultant left-shifted oxygen dissociation curve. (For this test, contact Special Haematology, St Thomas', London on 020 7188 3421 or Department of Haematology, City Hospital, Birmingham, UK, on 0121 507 4577 to discuss. A 5 ml EDTA sample will be required from the patient and a normal control, bled at the same time.)

**Erythropoietin receptor and von Hippel–Lindau (VHL) gene mutation analysis**

Patients with an unexplained erythrocytosis and low serum EPO levels should be considered for investigation of an EPO receptor mutation (Percy et al., 1998). The Chuvash form of erythrocytosis, an autosomal recessive disorder common to a large number of families in central Russia, has been shown to result from mutations in the VHL gene. Recently, a small number of patients of Asian and Western European ancestry with erythrocytosis have also been reported to have VHL mutations (Percy et al., 2003). These patients have inappropriately normal or high EPO levels for their Hct. (Investigation of EPOR and VHL gene mutations can be analysed by sending an EDTA sample by post to Dr M Percy, Department of Haematology, C Floor, Belfast City Hospital, Lisburn Road, Belfast, UK, 028 9032 9241, ext. 3325.)

**PRV-1 and other investigations**

A novel cell surface receptor Polycythaemia rubra vera-1 (PRV-1) has been found to be overexpressed in granulocytes from PV patients. Decreased c-MPL protein expression has been found in platelets from patients with PV. These tests are interesting but not currently useful diagnostic tests (Kralovics et al., 2003).

**Polycythaemia Vera**

Polycythaemia vera presents at a median age of 60 years. The incidence is the same in males and females. Reports of the annual incidence of PV vary widely, from 0.2/106/year in Japan (Kurita, 1974) to 28/106/year in Goteborg, Sweden (Kutti & Ridell, 2001). Patients often present with either arterial or venous vascular occlusive events (Barabas et al., 1973). Coronary and cerebral events are prominent and microvascular disturbances can also occur (Gruppo Italiano Studio Policitemia, 1995). Occasionally, they can present with haemorrhage particularly involving the skin and gastrointestinal tract. Splenic pain, pruritus, gout and constitutional symptoms, such as fatigue, may also be presenting features.

The disease may progress over time in various ways. Thrombosis and haemorrhage continue to occur. Splenomegaly may develop and increase. A few patients will proceed to develop massive splenomegaly and some myelofibrosis but there is variation in the terminology used. Older terms such as ‘spent phase’, ‘burnt out PV’, and ‘postpolycythaemic myeloid metaplasia’ are poorly elucidated. Modern definitions have relied on the demonstration of significantly increased bone marrow reticulin staining (grade III–IV on a four-point scale) together with a clinical syndrome of one or more of progressive splenomegaly, anaemia, leucoerythroblastosis and constitutional symptoms (Thiele et al., 2001b). With this definition, the rates of myelofibrotic transformation range from <5% (Berk et al., 1995; Tatarsky & Sharon, 1997), up to 15% (Najean & Rain, 1997a) at 10 years. Acute leukaemia is part of the natural history of PV, occurring in untreated patients and in those treated with venesection only (Berk et al., 1986). Historically, untreated PV had a dismal prognosis as shown by an early retrospective cohort study (Chievitz & Thiede, 1962), where the median survival in untreated patients was 18 months, with thrombosis being the dominant cause of death.

The aims of treatment of PV are therefore to:

I. reduce the risk of thrombosis and haemorrhage;
2 minimise the risk of transformation to acute leukaemia and myelofibrosis;
3 manage complications which may occur including thrombosis, haemorrhage and pruritus; and
4 manage pregnancy.

These aims were used when assessing the evidence and developing management advice.

**Management of polycythaemia vera**

*Randomised clinical trials*

There have been six randomised clinical trials of treatment for PV (Table V). Most trials have used the PVSG definition for the diagnosis of PV as the inclusion criterion, ensuring that the study populations are broadly comparable. None of the trials are without methodological problems. Most of these arise in studies of any disease with a long, indolent course, aging patient populations and significant late complications. It also proved difficult to keep patients on allocated treatment when new information became available. No trial had an objective of controlling the platelet count. They have also been reported in different ways. The issue of incidence of leukaemia is particularly problematic, as in some cases the actual incidence of acute leukaemia of a group of patients on a particular treatment is reported and in others the calculated actuarial risk on an intention to treat basis is reported. These figures are not comparable. However, there are many clear conclusions about the short and longer-term consequences of various therapeutic options that can be reached from review of the clinical trial data.

The PVSG-01 study established venesection as the first line therapy for PV. In comparison with $^{32}$P and chlorambucil, overall survival was significantly longer in the venesection arm, and associated with much lower risks of leukaemia and non-haematological malignancy. An increased risk of thrombosis was seen in the venesection arm, but this was predominantly observed during the first 3 years when the target Hct was 0Æ52 (Berk et al, 1995). In later years, it was reduced to 0Æ45. There was a large degree of cross-over between arms, with 91% of patients randomised to venesection having changed to alter-

<table>
<thead>
<tr>
<th>Table V. Randomised trials in polycythaemia vera with rates of important end-points for each of the treatment arms.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trial</td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td>PVSG-01§</td>
</tr>
<tr>
<td>$^{32}$P</td>
</tr>
<tr>
<td>Chl</td>
</tr>
<tr>
<td>PVSG-05¶</td>
</tr>
<tr>
<td>$^{32}$P</td>
</tr>
<tr>
<td>EORTC**</td>
</tr>
<tr>
<td>$^{32}$P</td>
</tr>
<tr>
<td>FPSG &gt;65 years††</td>
</tr>
<tr>
<td>$^{32}$P + HU</td>
</tr>
<tr>
<td>FPSG &lt;65 years‡‡</td>
</tr>
<tr>
<td>Pipob</td>
</tr>
<tr>
<td>ECLAP§§</td>
</tr>
<tr>
<td>Placebo</td>
</tr>
</tbody>
</table>

*Variables which are significantly different between arms in the trial.
†Estimates which have been derived from actuarial survival curves for the purposes of this table.
‡selected patients, not intention to treat.
¶Tartaglia et al (1986).
††Najean & Rain (1997a).
‡‡Najean & Rain (1997b).
native treatments by 10 years in the French subgroup of patients in the trial (Najean et al., 1994). Thus the role of purely using venesection as treatment for PV is unclear.

PVSG-05 was a two-arm study that compared $^{32}$P with venesection plus high doses of anti-platelet agents, aspirin (300 mg t.i.d.) and dipyridamole (75 mg t.i.d.) (Tartaglia et al., 1986). The rationale for this trial was to use anti-platelet agents to reduce the increased risk of thrombosis that was observed initially in the phlebotomy arm of PVSG-01. The haemorrhage and death rate was significantly increased in the venesection and antiplatelet arm and the trial was therefore stopped. In the majority of cases a high platelet count was found at the time of haemorrhage but the platelet count was controlled by $^{32}$P in the other arm.

The European Organisation for Research on Treatment of Cancer (EORTC) conducted a trial comparing $^{32}$P to busulphan (EORTC, 1981). Venesection was added in each arm to maintain the Hct at between 0.42 and 0.47. Overall survival was significantly better in the busulphan group, with the major reason for the difference being an increase in vascular complications. There were no differences between the arms for other complications, such as leukaemia, myelofibrosis or non-haematological malignancy.

The French Polycythaeemia Study Group (FPSG) published two randomised trials in 1997. The first was a two-arm comparison of $^{32}$P alone against $^{32}$P with maintenance hydroxycarbamide (formerly known as hydroxyurea) in patients over the age of 65 years (Najean & Rain, 1997a). Significant numbers of patients crossed between the two treatment arms. Median survival was not significantly different. No differences were observed for vascular end-points or progression to myelofibrosis. The actuarial risk of leukaemia was significantly greater for the $^{32}$P and hydroxycarbamide group, with the difference becoming apparent after 5 years, and the gap continuing to widen up to the 15th year. In addition, the actuarial risk of non-haematological malignancy was also much greater for the $^{32}$P and hydroxycarbamide arm, with a similar 5–15 year latency observed.

The second trial from the FPSG was a comparison of hydroxycarbamide therapy with pipobroman in patients under the age of 65 years (Najean & Rain, 1997b). Overall actuarial survival was 70% in the two arms at 14 years, compared with an estimated 84% for the age- and sex-matched population. There were no differences between the two groups in vascular end-points or rates of leukaemia or non-haematological malignancy. Myelofibrosis risk was significantly increased in the hydroxycarbamide arm, and tended to occur earlier.

The FPSG (Najean et al., 1994) also document very long follow-up in their patients entered into PVSG trials. They had 48 patients in PVSG-01 on chlorambucil, 60 in PVSG-01 and 21 in PVSG-05 on $^{32}$P, 56 in PVSG -01 and 19 in PVSG-05 managed with venesection. Of the original 75 patients randomised to venesection, 91% were treated with chemotherapy or radiotherapy. The numbers are thus so small that any analysis of survival is of very dubious accuracy. However, the patients in the phlebotomy arm developed myelofibrosis earlier but had very long survival. Therefore, intention to treat analysis is not valid. The risk of myelofibrosis was higher if treated by phlebotomy alone.

The European Collaboration on Low-dose Aspirin in Polycythaemia vera (ECLAP) study established the therapeutic benefit of aspirin in PV (Landolfi et al., 2004), and followed an earlier pilot study (Gruppo Italiano Studio Policitemia, 1997). Patients were randomised between aspirin 100 mg/d and placebo. Aspirin significantly reduced the risk of the combined end-point of non-fatal thromboembolic events, or death from cardiovascular causes. The risk of major or minor thrombosis was also significantly decreased. There was no significant increase in haemorrhage. The results of this large, well-designed multicentre trial eliminated the concerns about the efficacy and safety of aspirin that were raised by the earlier, smaller PVSG-05 trial (Tartaglia et al., 1986) and provided evidence for the use of aspirin in the management of PV.

The role of the platelet count

Hyperviscosity secondary to the raised Hct is a well-recognised cause of thrombosis in PV. There is a direct and striking correlation between the Hct and rates of thrombosis (Pearson & Wetherley-Mein, 1978). However, platelets may also be part of the problem. Circumstantial evidence for the benefit of reducing the platelet count is provided by studies in essential thrombocythaemia (ET) where reducing the platelet count significantly reduced the incidence of vascular events (Cortezlazzo et al., 1995). In the PVSG-01 trial, there was no correlation between a raised platelet count and thrombotic events (Berk et al., 1986). However, aspirin reduced the incidence of thrombosis in PV (Landolfi et al., 2004), suggesting a key role for platelets in thromboembolic events.

Different treatments vary little in the frequency of myelofibrotic transformation, although one randomised trial did show a greater incidence for hydroxycarbamide compared with pipobroman (Najean & Rain, 1997b). In this trial, poor control of the platelet count was strongly associated with progression to myelofibrosis. This suggests that controlling the platelet count will reduce thromboembolic events and influence the rate of transformation to myelofibrosis.

Venesection

An uncontrolled Hct has been associated with increased morbidity and mortality in surgical patients (Wasserman & Gilbert, 1964) and therefore ideally the Hct should be controlled for 3 months before elective surgery. Venesection can be used to control the Hct in PV. In a retrospective study, patients with PV who were treated with venesection and chemotherapy, mainly busulphan, the incidence of arterial and venous thromboembolic events increased as the Hct increased. The cerebral blood flow was significantly below normal in PV patients with a raised Hct and improved by 73% when the Hct
was less than 0·45 (Thomas et al., 1977). Thus, we recommend that the Hct is reduced to below 0·45 by venesection. There is currently no evidence to support a different level of Hct in males and females.

Isovolaemic erythropheresis has been used to reduce the RCM (Kaboth et al., 1997). To be used generally, it would have significant resource implications but it could be used acutely in the very occasional patient with an evolving ischaemic event.

**Recommendation: Venesection**

Venesection: the Hct should be maintained at less than 0·45 by venesection. The volume removed should be commensurate with the patient’s size and comorbidities.

Grade B recommendation: Evidence level IIa.

### Cytoreductive therapy

**Chlorambucil**

The alkylating agent chlorambucil was included in the randomised trial PVSG-01 (Berk et al., 1981) as listed in Table V. It was associated with a higher risk of acute leukaemia. The risk of acute leukaemia was higher with higher doses of chlorambucil. Chlorambucil is not now recommended in the treatment of PV.

**Busulphan**

The alkylating agent busulphan was used in the EORTC randomised trial and found to be superior to $^{32}$P (EORTC, 1981) (see Table V) although both treatments had a low incidence of acute leukaemia at a median follow-up of 8 years.

A single centre retrospective study (Messinezy et al., 1985) presented patients treated with venesection and low dose busulphan. The median survival was 11·1 years. Acute leukaemia and myelofibrosis deaths were significantly increased above the normal population but there were some long-term survivors with myelofibrosis. These studies show that low dose busulphan is efficacious in controlling PV but, since busulphan is an alkylating agent, it should be reserved for the elderly.

**Pipobroman**

Pipobroman is a bromide derivative of piperazine, similar to alkylating agents, which inhibits DNA and RNA polymerase and reduces incorporation of pyrimidine nucleotides into DNA. It has been used extensively in Europe but not in the UK. It has been used in a number of series of patients and one randomised trial (Najean & Rain, 1997b). Rates of acute leukaemia from 4% to 6% have been reported, except in one series where a rate of 19·5% was observed (Kiladjian et al., 2003) and in which some patients were also treated with other agents. Rates of myelofibrosis from 0% to 8·5% have been reported.

**Hydroxycarbamide**

Hydroxycarbamide acts by a non-alkylating mechanism by inhibiting the enzyme ribonucleotide reductase, which has a rate-limiting role in the regulation of DNA synthesis. It has been used in a phase II study, PVSG-08, and a number of case series. The PVSG-08 study was a Phase II efficacy study that included untreated and previously treated patients (Donovan et al., 1984). One-year failure-free survival was 73% in the previously untreated patients and 59% in the previously treated group. This study showed that hydroxycarbamide was efficacious but recommended dose reduction because of toxicity. The previously untreated patients in this study were compared with historical venesection randomised to venesection in PVSG-01 (Kaplan et al., 1986) with the Hct maintained at less than 50%. There was less thrombosis, including the early period after the start of therapy, and no difference in the rate of leukaemia. With prolonged follow-up, there was no statistical difference in the incidence of myelofibrosis (spent phase) between the groups (Fruchtman et al., 1997).

There has been much debate on the leukaemogenicity of hydroxycarbamide. In a series of reports on the use of hydroxycarbamide the incidence of acute leukaemia varied from 0% to 12% in patients treated with hydroxycarbamide alone (Weinfeld et al., 1994; Tatarsky & Sharon, 1997). The FPSG trial reported that patients aged over 65 years had an actuarial incidence of leukaemia of 12% in the hydroxycarbamide alone arm (Najean & Rain, 1997a). However, the FPSG randomized trial in patients below 65 years showed that pipobroman had a lower rate of progression to leukaemia and other cancers compared with the combination of $^{32}$P and hydroxycarbamide.

They also showed that progression to myelofibrosis occurred significantly more frequently on hydroxycarbamide in those over 65 years of age, but this was related to higher platelet counts in the hydroxycarbamide treated group.

The literature discussed above does not present conclusive evidence for increased leukaemogenicity of hydroxycarbamide given that the development of acute leukaemia is part of the natural history of PV. However, there is some related evidence that should be considered when evaluating hydroxycarbamide.

Over 10 years experience of the use of hydroxycarbamide in patients with sickle cell disease is now available and only four malignancies have been reported, two of which occurred soon after starting treatment and are probably not associated (Halsey & Roberts, 2003). However, caution should be applied in extrapolation of this to PV since sickle cell disease is not a clonal disorder. A study of patients with MPD exposed to hydroxycarbamide in vivo showed no increase in acquired DNA mutations compared with controls (Hanft et al., 2000). A series of patients with ET who developed 17p deletions following treatment with hydroxycarbamide (Sterkers et al., 1998) is often referred to as proof of the leukaemogenicity of hydroxycarbamide. The risk of leukaemia is this series was low (3·5%) in the patients treated with hydroxycarbamide alone.
The association with the particular cytogenetic abnormality was not proven in this group as there was no difference in the rate of 17p deletion in those treated with hydroxycarbamide alone and the rate in those not receiving cytotoxic agents. Therefore, the case for the leukaemogenicity of hydroxycarbamide is not proven. It is effective in controlling the counts and reducing the thromboembolic events.

Hydroxycarbamide is generally good at controlling PV. Continuous treatment is required and some patients will find this follow-up difficult. There will be some patients who will experience significant side effects, including gastrointestinal disturbance, skin pigmentation and leg ulcers. As there is still some anxiety about the possibility of leukaemia transformation its use should be limited in younger patients.

Radiotherapy and radioactive phosphorus $^{32}$P

The first patient was treated with $^{32}$P in 1940 (Lawrence, 1940). Within a few years cases of acute leukaemia were observed. A number of series of patients have been reported over the last 40 years. These series are often retrospective, of patients referred to tertiary centres and are likely to be selected cases that may have aggressive or problematic disease. It is of note that Osgood (1964) reported rates of acute leukaemia of 14% in PV patients and 2.5% in chronic lymphocytic leukaemia patients given the same treatment. This argues that acute leukaemia is part of the inherent nature of PV. The incidence in randomised trials after $^{32}$P alone is 2.5 to 15%.

$^{32}$P is good at controlling PV, intermittent treatment is required and follow-up can therefore be limited. However, it does increase the leukaemic transformation rate and therefore its use should be limited to the elderly.

Other cytotoxic agents

Melphalan, an alkylating agent, was used in one small series. Counts were well controlled but acute leukaemia developed in 15% and myelofibrosis in 14% of cases (Logue et al, 1970). The antimitabolite 6-thioguanine, which is a purine antagonist analogue of guanine, was used in one small, single-centre series and was of some efficacy (Milligan et al, 1982). The alkylating agent, carboquone and other agents were given in a single centre series (Higuchi et al, 1995). High rates of acute leukaemia resulted.

Interferon-α

Interferon-α (IFN-α) suppresses the proliferation of both pluripotent and lineage-committed haematopoietic progenitors. In vivo an inhibitory effect on progenitor cells is consistent with suppression of proliferation as the major mechanism controlling thrombocytosis and erythrocytosis. There are a number of small, single-centre series of patients treated with IFN-α for the control of PV. Complete response rates between 29% and 86% have been reported (Taylor et al, 1996; Foa et al, 1998; Heis et al, 1999). The definition of response varies, but IFN-α controls the blood counts, reduces or obviates the need for venesection and often reduces the amount of splenomegaly. It is also effective in many cases in reducing symptomatic pruritus. This agent has not been implicated as a possible leukaemogenic agent. However, it is not well tolerated in many patients as shown by withdrawal rates of up to 41%. One study (Heis et al, 1999) also documented a reduced rate of venous thrombosis on treatment but not the rate of arterial thrombosis compared with the rates on prior treatments.

IFN-α is theoretically superior for treating PV as it is effective in controlling counts and there is no risk of leukaemogenesis. Treatment is usually continuous but occasionally can be stopped for prolonged periods of time. The rate of side effects may make it difficult to tolerate. It is most likely to be tolerated in younger patients for whom it is recommended.

Busulphan, hydroxycarbamide and IFN-α can all reduce slight to modest splenomegaly but none are particularly efficacious in reducing massive splenomegaly.

Anagrelide

Anagrelide is a quinazolon derivative that inhibits cyclic nucleotide phosphodiesterase and the release of arachidonic acid from phospholipase, possibly by inhibiting phospholipase A$_2$. It was shown to control the platelet count in MPD patients including PV and side effects included cardiac, gastrointestinal and neurological. At 5 years, 16% of patients had discontinued this treatment because of side effects (Anagrelide Study Group, 1992). The results of the current UK Medical Research Council PT-1 trial may provide more information on the efficacy of anagrelide. Anagrelide is megakaryocyte-specific. It is effective in controlling the platelet count but probably does not control progression of PV, e.g. the increasing splenomegaly. The side effect profile may make it difficult for some patients to tolerate.

Recommendations

Drawing together the evidence available from randomised trials and other evidence we make the following recommendations for the management of PV.

Recommendations: Management of polycythaemia vera.

- Venesection to maintain the Hct to <0.45.
- Aspirin 75 mg/d unless contraindicated.
- Cytoreduction should be considered if:
  - poor tolerance of venesection;
  - symptomatic or progressive splenomegaly;
  - other evidence of disease progression, e.g. weight loss, night sweats;
  - thrombocytosis.
Thrombotic and haemorrhagic complications

Thrombotic risk assessment

Thrombotic events are a major cause of morbidity and mortality in PV. Clearly part of this risk relates to derangement of the full blood count (FBC), principally the Hct and platelet count but other factors may also be important. Conventional risk factors for atherosclerosis, including hyperlipidaemia and hypertension, have been assessed in MPDs with variable results. Little work has specifically been performed in PV. Recent recommendations for the management of atherosclerosis would suggest that this patient group would benefit from aggressive risk management with the use of antihypertensives to maintain normal blood pressure and the use of a statin. The utility of inherited thrombophilia screening in patients without MPD and with previous venous thromboembolism (VTE) has recently been questioned (Baglin et al, 2003). A recent report suggests that Factor V Leiden may be more common in MPD patients with recurrent VTE (Ruggeri et al, 2002). Although hyperhomocysteinaemia has been documented in patients with ET and PV, this appears neither to be associated with thrombosis nor the \( \text{MTHFR} \) polymorphism (Faurusch et al, 2000). Thus, there is no evidence as yet that identification of an inherited thrombophilic abnormality adds to the management of patients with MPD. Clinical studies are required to resolve this issue. An increased prevalence of antiphospholipid syndrome (aPL) has been described in ET and associated with increased risk of thrombosis (Harrison, 2002). There have been no reports thus far in this field for patients with PV. Patients with persistent antiphospholipid antibodies should be managed according to guidelines for this condition (Greaves et al, 2000).

Recommendations: Assessing risk of thrombosis

- Patients should be screened for hypertension, hyperlipidaemia, diabetes and a smoking history taken.
- Conventional risk factors for atherosclerosis should be managed aggressively. All patients should be requested to stop smoking.

No current evidence to support routine thrombophilia screening in PV.

Grade C recommendation: Evidence level IV

Management of an acute thrombotic event and pharmacoprophylaxis

Acute thrombotic events should be managed according to current guidelines, individual risk factors should be examined and control of the Hct and platelet count optimised. Low-dose aspirin has clear benefit in the secondary prevention of atherosclerosis in haematologically normal patients. However, its role in preventing similar complications for patients with MPDs has been controversial with the documented ability of aspirin to uncover a latent bleeding tendency (Tartaglia et al, 1986). Two subsequent studies of low dose aspirin (100 mg/d) in ET documented its safety (if platelet count <1000 x 10^9/l and no prior haemorrhage) as well as potential ability to reduce arterial thrombotic complications (van Genderen et al, 1995). As discussed, the recent ECLAP study (Landolfi et al, 2004) supports the safety and utility of aspirin in the prevention of non-fatal thrombotic events in PV. Unlike aspirin, current risk benefit analysis would suggest no role for oral anticoagulants, such as warfarin, in the primary prevention of thrombosis. For secondary prevention of VTE it remains unclear whether to give a short course (standard practice) or to continue with long-term warfarinisation. The only clear indication in this context is if the patients have antiphospholipid syndrome.

The role of the ADP-receptor antagonist clopidogrel in MPD patients is currently unclear. In patients with established atherosclerosis who do not have MPD, the recent CURE study (Clopidogrel in Unstable Angina to Prevent Recurrent Events) showed that combination therapy produced a 20% relative risk reduction of cardiovascular events and death, but was associated with a relative increased risk of major bleeding events of 38% (Yusuf et al, 2001). This must be of concern in the MPD patient population. Evidence exists for its benefit in patients with ongoing evidence of platelet activation whilst receiving aspirin (Nurden et al, 1996). These agents may be useful for patients with peptic ulcer disease or aspirin allergy.

Haemorrhage

Haemorrhage is both a less frequent and generally less severe clinical complication of PV than thrombosis. The principal sites affected are skin, mucous membranes and gastrointestinal tract. Haemorrhage is often reported in association with high platelet counts, acquired von Willebrand disease (Budde et al, 1984) and high doses of anti-platelet therapy (Tartaglia et al, 1986). Low dose aspirin is infrequently associated with haemorrhagic complications (Landolfi et al, 2004). A wide
Guideline

variety of platelet function defects are reported in PV but they are not predictive of bleeding.

Clinically significant bleeding may paradoxically require platelet transfusion (Terasako & Sasai, 1998) and a role for epsilon amino caproic acid and tranexamic acid has been suggested by some (Spivak, 2002). Other measures to consider in the uncommon, less acute situation include better control of blood counts, adjustment of any concomitant anti-platelet and/or anti-coagulant therapy. The utility of recombinant Factor VIIa is unknown in MPD patients with uncontrolled life-threatening bleeding and is worthy of further study.

Pruritus

Pruritus, typically aquagenic, can be a severe clinical problem in PV. Antihistamines may be of benefit (Weick et al, 1982). A number of small studies have tried to address this problem. Some benefit has been shown with cimetidine (Weick et al, 1982) and phototherapy using psoralen and ultraviolet A light (Jeanmougin et al, 1996). One study showed improvement in pruritus with iron replacement but the pruritus recurred when iron had to be stopped (Hct controlled to 0.55) (Salem et al, 1982). Some studies (Taylor et al, 1996) described improvements with treatment with IFN-α. Finally, Tefferi and Fonseca (2002) reported 10 patients with PV who were treated with selective serotonin re-uptake inhibitors for other reasons and showed great improvement in their pruritus. There are thus a number of agents that may be useful on an individual case basis.

Pregnancy and polycythaemia vera

There is only limited information in the medical literature about the management of PV in pregnancy. Of the 20 pregnancies reported there were 12 live births but three of 12 suffered early neonatal death; 17 of 20 cases were reported before 1988. A recent series of 20 pregnancies (Robinson et al, 2004) reviewed a further 16 pregnancies in eight women and documented significantly greater chance of live birth with aggressive management. The risks of pregnancy in PV are probably similar to those for patients with ET where there is a more extensive literature with an overall incidence of first trimester miscarriage of 36% (about twice that expected) and an increased chance of intrauterine growth retardation, intrauterine death and stillbirth described (8%) (Harrison, 2002).

An overview of the literature does not enable confident management guidelines to be drawn up. These recommendations are based on current knowledge of PV, ET and the management of antiphospholipid syndrome, which all have placental dysfunction as a common pathogenic feature. Therapeutic strategies for PV in pregnancy are influenced by the patients’ disease status and prior obstetric history. If any of the following factors are present, then the pregnancy is likely to be at high risk of complication to the mother and/or fetus:

- previous venous or arterial thrombosis in mother (whether pregnant or not);
- previous haemorrhage attributed to PV (whether pregnant or not);
- previous pregnancy complication that may have been caused by PV; e.g. ≥3 first trimester or ≥1 s or third trimester pregnancy loss; birthweight <5th centile for gestation;
- intrauterine death or still birth (with no obvious other cause, evidence of placental dysfunction and growth restricted fetus);
- significant ante- or postpartum haemorrhage (requiring red cell transfusion);
- severe pre-eclampsia (necessitating preterm delivery <37 weeks) or development of any such complication in the index pregnancy;
- platelet count rising to >1000 × 10^9/l.

Therapeutic options include antithrombotic treatment, venesection and cytoreductive agents, although the expected natural fall of the platelet count and Hct during pregnancy may anyway obviate or reduce the need for the latter. The Hct could be controlled with either careful venesection or cytoreductive therapy. The target Hct for a non-pregnant female has yet to be determined, but in pregnancy the Hct should be maintained within the normal range appropriate for gestation. There is currently no evidence for maintaining Hct less than this in pregnancy.

Cytoreduction should be avoided in pregnancy, particularly in the first trimester. None of the cytoreductive agents have a product licence for use in pregnancy. Where cytoreduction is deemed necessary (see above), IFN-α is the drug of choice. There are no reports of teratogenic effects in animals or adverse effects in the, admittedly, small numbers of pregnancies exposed to this drug. However, some evidence suggests that IFN-α may decrease fertility (Griesshammer et al, 1998) and so it may be better to avoid it in women who have difficulty in conceiving. Few pregnancies in chronic myeloid leukaemia patients treated with hydroxyurea have been published (Patel et al, 1991; Jackson et al, 1993), most without fetal complications. However, one still-birth and one malformed infant and teratogenicity in animals have been reported. Hence hydroxyurea is probably contraindicated at the time of conception (this also applies to male patients) and during pregnancy. Anagrelide is not recommended because of insufficient documentation of its use in pregnancy. Thus hydroxyurea or anagrelide should be gradually withdrawn 3–6 months prior to conception and IFN-α may be substituted if necessary.

Low dose aspirin is safe in pregnancy (Collaborative Low-dose Aspirin Study in Pregnancy (CLASP) Collaborative Group, 1994) and seems advantageous (Griesshammer et al, 1998). We recommend that, in the absence of clear contraindications, all patients should be on aspirin (initially 75 mg o.d.) throughout the pregnancy and for 6 weeks after delivery (grade C recommendation, evidence level IV).
Low molecular weight heparin (LMWH) has been used successfully in pregnancies at high risk of thrombosis (Hunt et al, 2003). It reduced fetal morbidity (Rai et al, 1997), is safe and has a lower risk of heparin-induced thrombocytopenia and osteoporosis compared with unfractionated heparin (Sanson et al, 1999). Thus it has been used anecdotally in women with PV and previous thrombosis and/or fetal morbidity. If the patient has had a previous venous or arterial thrombosis, then the use of LMWH thromboprophylaxis is indicated during pregnancy. Use of unmonitored intermediate dose LMWH is widely used (e.g. enoxaparin 40 mg o.d.) increased to 40 mg twice daily from 16 weeks, dropping to 40 mg/d for 6 weeks postpartum. The recent guidelines for thromboprophylaxis in pregnancy recommend constant reassessment of venous thrombotic risk during pregnancy and that all women with previous VTE or a thrombophilia should be encouraged to wear graded elastic compression stocking (GECs) throughout their pregnancy and for 6–12 weeks after delivery (Haemostasis and Thrombosis Task Force, British Committee for Standards in Haematology, 2001). The use of GECs is also recommended for pregnant women travelling by air (Kelman et al, 2003).

During the pregnancy the patient should be monitored regularly and management is summarised in Fig 1. It is important to discuss the implications of the use of thromboprophylaxis with the obstetric anaesthetist for epidural or spinal anaesthesia. During labour, dehydration should be avoided, attention should be given to the LMWH dose and the use of GECs should be considered. In the puerperium, we recommend thromboprophylaxis with 6 weeks LMWH for all women with MPD. Breast feeding is safe with heparin and warfarin (providing baby receives adequate vitamin K). Breast feeding is contra-indicated with the cytoreductive agents (IFN α, anagrelide and hydroxycarbamide). The first 6 weeks postpartum are a high risk time for venous thrombosis; blood counts may rise rapidly, thus on-going haematological monitoring is important.

**Apparent erythrocytosis**

Evidence that apparent erythrocytosis is associated with increased mortality comes from small, non-randomised studies in which the survival of patients with apparent erythrocytosis is compared with the expected death rate of age- and sex-matched controls in the general population. Burge et al (1975) studied 32 men and three women with a Hct of 0.50, RCM ≤36 ml/kg and a plasma volume ≤36 ml/kg. Twenty-seven of these patients were followed-up for a minimum of 4 years. There was a significant excess of deaths in this patient group over the expected death rate. Weinreb and Shih (1975) investigated 69 men with a RCM ≤36 ml/kg who were referred for evaluation of a Hct >0.52. Patients were sub-divided depending on whether their RCM was in the upper normal range (group 1) or was no greater than 1 SD from the normal mean (group 2). Patients in group 2 had a significantly lower plasma volume than those in group 1. The survival of patients in group 2 was poorer than in group 1 and group 2 patients had a poorer survival than age-matched controls in the general population.

Circumstantial evidence for an increase in morbidity and mortality in apparent erythrocytosis comes from studies showing that individuals with a Hct in the upper normal range, or slightly elevated, may be associated with an increase in thrombotic events and cardiovascular mortality compared with those with a Hct in the middle or lower part of the normal range (Lowe, 1999).

It is not clear that the increase in mortality associated with apparent erythrocytosis is because of the high Hct, nor are there randomised studies to show that reducing the Hct in apparent erythrocytosis reduces morbidity or mortality. Current management of apparent erythrocytosis may be based on the following observations.

Serial measurements of the Hct in untreated patients with apparent erythrocytosis show that the Hct returns to within the normal range in up to 30% of patients (Messinezy & Pearson, 1990).

Modifications in the factors associated with apparent erythrocytosis, such as obesity, smoking and hypertension, may lead to a reduction in Hct (Pearson, 1991).

Venesection of male patients with apparent erythrocytosis because of a low plasma volume resulted in a fall of mean Hct from 0.5 to 0.43, with a rise in plasma volume without a reduction in overall blood volume (Humphrey et al, 1980).

There is a need for more data on the clinical consequences of apparent erythrocytosis. If it is confirmed to be an independent risk factor for thrombosis, then randomised studies of treatment to lower the Hct are required, on which to base rational management.

**Recommendations: Management of apparent erythrocytosis**

Confirm that the elevated Hct is persistent, with at least two measurements of the Hct under standardised conditions over a 3-month period.

Advise reduction or elimination of factors which may contribute to apparent erythrocytosis, e.g. a reduction in smoking and alcohol intake and control of hypertension (without the use of a thiazide diuretic).

Consider venesection in the following circumstances:

- Patients with a recent history of thrombosis, or with additional risk factors for thrombosis.
- Patients whose Hct exceeds 0.54 (>3 standard deviations above the mean), based on the increased risk of thrombosis in idiopathic erythrocytosis and low incidence of normal individuals with a Hct >0.54.
Summary of pregnancy management and LMWH doses

- FBC every 4 weeks until 24 weeks
- Then 2 weekly FBC
- BP + urinalysis every visit
- USS Scan
  - At 12, 20, 26, 30, 34 and 38 weeks
- Uterine Artery Dopplers
  - At 20 (+24 weeks if abnormal)

**Normal**

- Treat as normal

**Abnormal i.e. bilateral high RI or notches**

- Increase intensity of monitoring
- Consider:
  - Escalating LMWH dose
  - Add Vitamin C 1000 mg od
  - Vitamin E 400 iu od
  - Early delivery prior to 38 weeks

Follow local guidelines regarding anaesthetics and aspirin/LMWH

**Delivery**

- Stop LMWH once the patient goes into labour
- For elective caesarian section omit from 12 hours pre-procedure
- Follow local guidelines for regional/epidural anaesthesia
- Restart LMWH ASAP post-partum providing no bleeding

**Post partum**

- Continue aspirin ± LMWH for at least 6 weeks post-partum
- Control of maternal platelet count and PCV as normal

**Breast feeding** is contraindicated if a patient is having any cytoreductive therapy

**Low molecular weight heparin doses**

- Start when pregnancy test is positive, give subcutaneously

  **If normal body weight, no renal impairment + previous venous thrombosis or fetal morbidity**
  - Dalteparin 5000 in or enoxaparin 40 mg once daily
  - At 16-20/40 increase to twice daily
  - 3 days post-partum reduce to dialy for 6 weeks

  **If previous arterial event**
  - Dalteparin 5000 in or enoxaparin 40 mg twice daily throughout pregnancy
  - If evidence of recurrence consider increased LMW heparin dose or warfarin after 14/40

Fig 1. Summary of pregnancy management and LMWH doses.
Untreated patients should be monitored to exclude a further rise in Hct and possible evolution to absolute erythrocytosis.

There is no data on which to base a target Hct for patients undergoing venesection, but a Hct <0.45 has been proposed based on data from patients with PV and idiopathic erythrocytosis (Pearson, 1991).

Grade C recommendation: Evidence level IV.

**Idiopathic erythrocytosis**

The term idiopathic erythrocytosis applies to patients who have an increased RCM and who, on investigation, do not have any form of known primary or secondary erythrocytosis. It has also been termed ‘benign erythrocytosis’ but these patients do not always have a benign course (Modan & Modan, 1968) and ‘pure erythrocytosis’, as it was thought to be a pure red cell disorder (Najean et al, 1981). However, secondary erythrocytosis is also a pure red cell disorder, therefore, idiopathic erythrocytosis is the preferred term. There is a male preponderance in several of the published series (Modan & Modan, 1968; Pearson & Wetherley-Mein, 1979). The incidence of vascular complications is high, 46-6% at presentation in one series and 17% of the total patients died of cerebrovascular accidents (Pearson & Wetherley-Mein, 1979). In another study, the incidence of fatal thromboembolic and haemorrhagic events was the same as in patients with PV (Modan & Modan, 1968). A management plan must take this into account.

**Recommendations: Idiopathic erythrocytosis**

- Venesection to reduce the Hct to <0.45 if Hct >0.54.
- Venesection to reduce the Hct to <0.45 if <0.54 and there is increased risk of thrombosis, i.e. evidence of ischaemia, previous history of thrombosis, peripheral vascular disease, diabetes or hypertension.
- Cytoreductive therapy is contraindicated.

Grade B recommendation: Evidence level III.

**High oxygen affinity haemoglobins**

The physiological adaptations to the inheritance of a high oxygen affinity haemoglobin include a rise in Hct, which is often modest, and an increase in cardiac output. Healthy individuals with a high oxygen affinity haemoglobin and comparable p50 values have differing haemoglobin concentrations, suggesting heterogeneity in the adaptive responses (Charache et al, 1978). There have been no randomised studies on venesection therapy for patients with high oxygen affinity haemoglobins, and proof of the efficacy of this treatment is lacking.

Recommendations for venesection are based on, and influenced by, the following observations:

1. Most individuals remain asymptomatic.
2. Experimentally, isovolaemic venesections performed in asymptomatic individuals with a high oxygen affinity haemoglobin, and resulting in a reduction in Hct from mean values of 0.55 to 0.41, can reduce exercise performance (Butler et al, 1982; Winslow et al, 1983).
3. Hyperviscosity symptoms and thromboembolic episodes have been reported (Fairbanks et al, 1971; Weatherall et al, 1977).
4. In some families, thrombotic episodes have been confined to individuals who are compound heterozygotes for both a high oxygen affinity haemoglobin and a thrombophilic defect (Berruyer et al, 1994; Hanss et al, 2002).
5. Individuals with dizziness, dyspnoea or angina may derive clinical benefit from venesection (Fairbanks et al, 1971; Grace et al, 1992).

**Recommendations: High oxygen affinity haemoglobins**

Possible indications for venesection include the following:

- Presence of symptoms such as dizziness, dyspnoea or angina, for which a raised Hct is considered to be a contributory factor.
- One or more previous thrombotic episodes.
- Asymptomatic individuals in whom a family member with a high oxygen affinity haemoglobin, similar haemoglobin concentration, and comparable risk factors for thrombosis, has developed thrombotic problems.

Consideration of a partial exchange transfusion should be given for individuals with a Hct >0.60 requiring major surgery (Larson et al, 1997). Do not attempt to reduce the Hct to within the normal range. Venesection to maintain the Hct <0.60 has been recommended (Weatherall et al, 1977).

When thrombosis or symptoms compatible with hyperviscosity develop at a lower Hct, a target Hct of 0.52 has been suggested (Pearson T, personal communication).

Grade C recommendation: Evidence level IV.

**Hypoxia**

Erythrocytosis secondary to hypoxia occurs in different ambient circulatory conditions from the other forms of erythrocytosis. This influences the compensatory mechanisms for blood flow and also impacts upon management of the erythrocytosis. The challenge in managing these patients is balancing oxygen transport against the effects of increased viscosity because of the elevated Hct. Two areas will be
considered in detail: hypoxic pulmonary disease (HPD) and cyanotic congenital heart disease (CCHD).

**Hypoxic pulmonary disease**

The development of an erythrocytosis in patients with HPD is associated with an increased risk of the development of cor pulmonale and a poor median survival of 2–3 years (Criner, 2000). Additional factors affecting circulatory compromise and tissue oxygen delivery include carbon monoxide in smokers, extent of hypercapnia, renal blood flow, acid–base balance (pH), capacity of the bone marrow to respond to erythropoietic drive, position of the oxygen dissociation curve and changes in the peripheral vascular circulation (Harrison & Stokes, 1982). Furthermore, for an individual patient, co-existent age-related vascular disease may also affect therapy.

Long-term oxygen therapy improves survival in patients with chronic obstructive pulmonary disease and severe hypoxaemia (PaO₂ <7.3 kPa or <8.0 kPa with nocturnal hypoventilation) (Crockett et al., 2001). This also reduces the Hct by improving oxygenation. All patients with erythrocytosis consequent upon HPD should be evaluated by a respiratory physician for consideration of long-term oxygen therapy or alternative methods of improving oxygenation. If they are smokers they should be strongly advised to stop. In addition to supplemental oxygen, nocturnal oxygenation may also be improved by the use of non-invasive ventilation in the case of Type II respiratory failure (Simonds & Elliott, 1995) or continuous positive airways pressure, in the case of obstructive sleep apnoea (Jenkinson et al., 1999). Therefore, failure to achieve adequate oxygenation in HPD should not be accepted without review by a specialist respiratory physician. In addition, a clinically relevant minority of patients with erythrocytosis have nocturnal oxygen desaturation because of obstructive sleep apnoea (Moore-Gillon et al., 1986) and such patients should be referred for appropriate investigation (Eisensehr & Noachtar, 2001).

Benefit of limited venesection in patients with HPD was demonstrated by Weisse et al. (1975), who showed that reducing the Hct to 0.50–0.52 led to an improvement in exercise tolerance, but a further staged reduction to Hct of 0.45 did not give additional benefit. Numerous other non-controlled patient series have also suggested that control of the Hct reduces pulmonary vascular resistance (Segel & Bishop, 1966; Harrison et al., 1973; Weisse et al., 1975; Harrison & Stokes, 1982), improving cerebral blood flow and psychometric testing (Menon et al., 1981; Wedzicha et al., 1983) as well as subjectively helping confusion and headache (Wade et al., 1981). Not all studies, however, have demonstrated beneficial effects of venesection in these patients (Dayton et al., 1975) and there are reports of venesection fatalities (Constantinidis, 1979). It is therefore of interest that a number of other agents have been reported to reduce the Hct in small, uncontrolled studies. These include dapsone and pyrimethamine (Pengelly, 1966) as well as theophyllines (Oren et al., 1997) and losartan (Vlahakos et al., 1999).

**Recommendations: Hypoxic pulmonary disease**

- Patients with HPD who develop an erythrocytosis should be evaluated by a respiratory physician for consideration of long-term oxygen therapy or alternative therapy (Grade A recommendation: Evidence level IA).
- Patients who are symptomatic of hyperviscosity or have a Hct >0.56 should have venesection to reduce this to 0.50–0.52 (Grade B recommendation: Evidence level III).
- There is limited evidence to suggest that therapy with drugs such as angiotensin-converting enzyme (ACE) inhibitors or angiotensin receptor antagonists could be used in patients who do not tolerate venesection (Grade B recommendation: Evidence level Ila)

**Cyanotic congenital heart disease**

In CCHD, a compensatory erythrocytosis develops to maintain tissue oxygen delivery. The underlying heart defects fall into two large categories. (i) Patients with absent or poorly developed central pulmonary arteries with pulmonary blood flow via collateral arteries from the aorta or branches and a large right to left shunt (typically pulmonary atresia with a ventricular septal defect and major aorto-pulmonary collateral arteries). (ii) Patients with pulmonary vascular disease where the intrapulmonary arterioles and capillaries have undergone obliterative changes secondary to high pressure and high flow shunts in a variety of simple and complex congenital heart defects and, occasionally, as a result of palliative surgical shunting procedures (Eisenmenger syndrome). Many of these patients are currently seen only in adult cardiology or haematology departments (Rosenthal & Anderson, 1998) who might now be eligible for corrective or palliative surgery or catheter interventions to decrease the cyanosis and the erythrocytosis.

As the erythrocytosis increases, patients may experience symptoms of hyperviscosity, though many may remain free from symptoms for many years, even with a Hct >0.70 (Perloff et al., 1988).

Thrombosis has been documented in several series of patients with CCHD. Interestingly, children, to whom this guideline is not intended to apply, appear to have a significantly increased risk of cerebral venous thrombosis that is particularly linked to iron deficiency (Phornphutkul et al., 1973). In adults, however, there is evidence of the occurrence of cerebral arterial and microarterial thrombi and particularly in patients with Fallots’ tetralogy pulmonary thrombotic events (Rich, 1948; Berthrong & Sabiston, 1951). Direct evidence linking thrombus to increased viscosity, however, remains at best circumstantial, in that older patients with cerebral thrombosis tend to be those with the highest Hct.
values (Phornphutkul et al., 1973). A similar link was not been shown in other case series (Perloff et al., 1993) but the series was probably not sufficiently powered to detect a difference (Bridges, 1994). A statistically significant correlation was also suggested between stroke, microcytosis and history of phlebotomy in one study of 162 adults with CCHD (Ammash & Warnes, 1996). It was not clear from this study when the stroke occurred in relationship to the venesection. There has also been a report of myocardial ischaemia directly related to high Hct in these patients (Yeager & Freed, 1984).

Cyanotic congenital heart disease patients who experience symptoms of hyperviscosity (Table VI) respond with a reduction in these symptoms with venesection. Venesection also results in reduced peripheral vascular resistance, and increased stroke volume, cardiac output and systemic blood flow (Rosenthal et al., 1970; Oldershaw & Sutton, 1980). Repeated exercise tests 2 weeks after venesection in one study showed increased oxygen uptake and reduced oxygen debt (Oldershaw & Sutton, 1980). The clinical difficulty arises when discriminating hyperviscosity from heart disease-related symptomology, and a pitfall to be avoided is deciding whether to venesection based upon the Hct alone. Furthermore, excessive venesection renders these patients both hypoxic and anaemic, reducing tissue oxygenation and exercise tolerance (Somerville, 1997).

There have also been concerns that, when venesection renders patients iron deficient, it may further increase blood viscosity and increase the risk of thrombosis (Hutton, 1979; Linderkamp et al., 1979). There are several pitfalls in this field; inaccurate Hct estimation in the presence of microcytic indices, and conflicting evidence as to whether iron deficient red cells are more rigid than normal red blood cells. On balance, the literature would suggest that iron-deficient red cells are at least as deformable as normal red cells (Pearson, 2001). However, a key problem when iron deficiency develops in CCHD is that the oxygen carrying capacity is no longer optimum (Gidding & Stockman, 1988). For example, at an MCH of 20 pg and Hct of 0.50, the haemoglobin and hence the oxygen carrying capacity are 11% less than for normal red cells (Van de Pette et al., 1986). The corollary is that iron-deficient patients with hypochromic cells will have a higher Hct and increased viscosity (and symptoms) than iron-replete patients with a similar haemoglobin value and oxygen carrying capacity.

A significant feature of the erythrocytosis in patients with CCHD (when compared with PV) is the absence of a thrombocytosis – indeed, the platelet count is frequently low – and evidence for platelet dysfunction. In addition, clotting factors II, V, VII and IX are reduced and may explain the mild bleeding diathesis in these patients (Territo & Rosove, 1991). These factors would seem to be theoretically advantageous in preventing stroke produced by the raised viscosity from the erythrocytosis. Indeed, it has been shown and is recommended that these patients should undergo a venesection 24 h before elective non-cardiac surgery to improve the clotting function (Wedemeyer & Lewis, 1973).

Thorne (1998) suggested isovolumic venesection should only be performed for symptomatic patients if the Hct is >0.65 and the patients had adequate iron stores. If the Hct is <0.65, iron deficiency should be suspected and, if present, this should be cautiously treated first as the hyperviscosity symptoms could be because of the iron deficiency itself (Perloff et al., 1993). Iron replacement, however, has been documented to cause a rapid rise in Hct or unstable erythrocytosis (Rosove et al., 1986) and so there are practical difficulties with these guidelines. A reasonable, evidence-based approach would be to venesection only symptomatic patients; if iron deficiency occurs then a small amount of iron could be administered with close supervision. Once the haematocrit begins to rise – often within a week – it should be stopped to prevent an exaggerated rise in the Hct. There are anecdotal reports of ischaemic symptoms with iron therapy and control of the Hct by venesection (Sondel et al., 1981).

In the absence of strong evidence associating the erythrocytosis with stroke development or evidence for a benefit from routine venesection in these patients, the development of symptoms from excessive venesection in some and the potential for stroke in those rendered iron-deficient, it seems reasonable for venesection to be restricted to those with symptoms of hyperviscosity. Antiplatelet agents and anticoagulation therapy for the prevention of stroke should be avoided given the increased bleeding tendency unless there are additional risk factors for stroke development, such as atrial fibrillation, poor ventricular function or a documented transient ischaemic attack. With continuing advances in surgical, catheter interventional and medical management of these patients, it would be reasonable to suggest that the management of CCHD patients is primarily in a specialist adult congenital heart disease unit with haematological support for the management of hyperviscosity symptoms (Table VI).

**Table VI. Symptoms of hyperviscosity in CCHD with erythrocytosis.**

<table>
<thead>
<tr>
<th>Symptom</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chest and abdominal pain</td>
</tr>
<tr>
<td>Myalgia and weakness</td>
</tr>
<tr>
<td>Fatigue</td>
</tr>
<tr>
<td>Headache</td>
</tr>
<tr>
<td>Blurred vision or symptoms to suggest amaurosis fugax</td>
</tr>
<tr>
<td>Paraesthesiae</td>
</tr>
<tr>
<td>Slow mentation, sense of depersonalisation</td>
</tr>
</tbody>
</table>

**Recommendations: Cyanotic congenital heart disease**

- Patients with CCHD and an erythrocytosis represent a complex management problem and should be managed primarily in a congenital heart disease unit so that advances in surgery, catheter interventional and medical management that may improve (rarely cure) the erythrocytosis are not missed. (Grade C recommendation: Evidence level IV)
Postrenal transplant erythrocytosis (PTE)

The correction of anaemia after a successful renal transplantation is dependent on the adequate production of EPO by the donor organ and the elimination of marrow inhibitors that characterise the uraemic state (Besarab et al., 1987). Following a successful transplant, the EPO production usually increases within a few days and results in a correction of the anaemia within 3 months. However, approximately 10–15% of renal transplant recipients develop an erythrocytosis between 8 and 24 months later, a condition referred to as postrenal transplant erythrocytosis (PTE) reviewed by Vlahakos et al. (2003). Pre-disposing factors include smoking, diabetes, transplant renal artery stenosis, rejection-free course with a well functioning renal graft and adequate erythropoiesis prior to transplantation. The pathogenesis of PTE is poorly understood and is likely to be multifactorial, involving abnormal erythroid precursor sensitivity to EPO or altered EPO production, abnormal erythroid sensitivity to angiotensin II or altered angiotensin II production and an elevated concentration of insulin-like growth factor I and its binding proteins (Gaston et al., 1994; Brox et al., 1998; Julian et al., 1998). PTE usually persists, with only 25% undergoing spontaneous remission, and presents clinically in most patients with malaise, headache, lethargy and dizziness. Importantly, PTE can contribute to the onset of hypertension or worsen pre-existing hypertension, and in addition, constitutes a serious thromboembolic risk factor. In one series, 10–30% of cases developed a thrombo-embolic event, that involved both veins and arteries, and which ultimately led to the patients’ death in 1–2% of cases (Wickre et al., 1983).

Management

It is important to prevent further increases in Hct levels by meticulous avoidance of extracellular volume reduction, for example excessive diuresis, diarrhoea and vomiting. PTE was originally treated with repeated venesection although this approach led to severe iron deficiency.

Recently, the beneficial effects of ACE inhibitors (ACEI) and angiotensin II receptor antagonists have been reported (Vlahakos et al., 2003). ACEI, including captopril, enalapril, lisinopril, are well tolerated and a dose-dependent decrease in Hct is seen within the first month of treatment with maximal effect being reached by 3 months. Similar results have been reported for the angiotensin II receptor antagonist, losartan. The minority of patients (5–0%) who fail to respond to one ACEI are not likely to respond to other ACEIs or to losartan, although an occasional patient may respond to a combination of an ACEI and theophylline (Rostaing et al., 1995). In the absence of response to these therapeutic interventions, venesection remains the only effective therapy and should be undertaken to achieve a target Hct of 0·45 (Barenbrock et al., 1993).

Recommendations: Postrenal transplant erythrocytosis

- Avoid excessive dehydration.
- Treat with ACEI or an angiotensin II receptor antagonist.
- Venesection to Hct of 0·45.

Grade C recommendation: Evidence level IV.

Useful web site http://www.acor.org/mpd/

Disclaimer

While the advice and information in these guidelines is believed to be true and accurate at the time of going to press, neither the authors, the British Society for Haematology nor the publishers accept any legal responsibility for the content of these guidelines.

Acknowledgements

We would like to thank Mrs Linda Megrath, Department of Haematology, Queen’s University, Belfast for her secretarial assistance in the preparation of this Guideline and Mr Dairmuid Kennedy, Librarian, Queen’s University, Belfast for help with the literature searches.

References


