Management of sickle cell disease: challenges and risks of transfusion

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Abstract: Homozygous sickle cell (SS) disease is associated with rapid red cell destruction and a tendency to block flow in blood vessels. The bone marrow expansion needed to compensate for the rapid red cell destruction increases metabolic demands and folate requirements but also renders the bone marrow prone to suppression by renal impairment and infections especially those with human parvovirus B19. The abnormal red cells also tend to block blood vessels impairing flow in the bone marrow (dactylitis, bone pain crisis, hip necrosis), the spleen (acute splenic sequestration, chronic hypersplenism, loss of the normal filtering system rendering patients prone to overwhelming septicemia), the lungs (pulmonary embolism, acute chest syndrome), and the brain (ischemic stroke, hemorrhage). Transfusion plays a role in addressing all of these pathologies. During the acute lowering of hemoglobin due to acute splenic sequestration and aplastic crisis and the persistent lowering of hemoglobin due to chronic hypersplenism and chronic renal failure, top-up transfusions may help in maintaining oxygen delivery and symptomatic relief. Addressing vaso-occlusion is more complex but best documented in preventing recurrent stroke and primary stroke following detection of cerebral vessel stenosis by transcranial Doppler. Although top-up transfusions have minimal side effects, potentially serious complications arise from chronic transfusion, and there are many unanswered questions on the duration of such therapy and the natural history of the underlying complications. These issues are addressed with the knowledge currently available.

Keywords: sickle hemoglobin, HbS, transfusion, oxygen affinity, anemia, vaso-occlusion

Introduction
The substitution of valine for glutamic acid at position 6 in the amino terminus of the beta chain in sickle hemoglobin (HbS) changes the behavior of the molecule. When deoxygenated, molecules of HbS tend to polymerize and deform the red cell resulting in the principal pathological features of the disease, hemolysis and obstruction of blood flow. The HbS mutations have occurred on at least 3 independent occasions in the African continent,¹ are named after the areas where they were first described, Benin, Bantu (or Central African Republic), and Senegal, and are known as the beta globin haplotypes. Most of the published knowledge on sickle cell disease has been derived from peoples of West African origin with the Benin haplotype. A fourth independent mutation occurred in the Arabian Gulf and Central India and is known as the Arab-Indian or the Asian haplotype.² This haplotype often has more mild clinical features because it is usually associated with high levels of fetal hemoglobin (HbF) and frequent deletional alpha thalassemia, both factors likely to inhibit intravascular...
Inheritance of the HbS gene from both parents result in homozygous sickle cell (SS) disease, which is generally severe and the most common at birth. Inheritance of the HbS gene along with another interacting hemoglobin variant results in doubly heterozygous conditions, sickle cell–hemoglobin C (SC) disease, sickle cell–beta thalassemia, and uncommon conditions such as sickle cell–HbO Arab, sickle cell–HbD Punjab, and sickle cell–hemoglobin Lepore. These conditions are generally milder than SS disease and often show differences in clinical and hematological expression. The current review is confined to SS disease but often distinguishes between African and Asian forms of the disease.

**Steady-state hematology**

The “steady state” refers to when a patient is clinically well and is characterized by hematological features, which oscillate within a narrow range in the same patient but can vary markedly between patients of the same genotype. In African SS disease, mean red cell survival is markedly shortened to 10–12 days, compared to 120 days in hematologically normal people, and requires expansion and increased turnover of the red cell precursors in the bone marrow. A new steady-state equilibrium (hemoglobin levels 6–9 g/dL, reticulocyte 10–12%, red mean cell volume (MCV) 80–95 fL) develops, knowledge of which is necessary to interpret the deviations from these values. Patients with the Asian haplotype SS disease may have different steady-state values (hemoglobin 8–11 g/dL, reticulocyte counts 3–8%, MCV 60–80 fL), consequent on the presumed greater red cell survival associated with high HbF and frequent alpha thalassemia common in this haplotype.

HbS within the red cell behaves with a low oxygen affinity with a marked shift in the oxygen dissociation curve to the right (Figure 1). Traditionally described as the p50 (oxygen tension associated with 50% saturation), values in normal AA blood are 29–31 mmHg, whereas in African SS disease, they may range from 40 to 60 mmHg. Patients with the greatest shift in p50 tend to have the lower steady-state hemoglobin levels, almost as if the erythropoietic drive is switched off at submaximal reticulocyte counts. This adaptation implies that hemoglobin level cannot be equated with oxygen-carrying capacity, and it is vital to treat the patient rather than the hemoglobin level.

**The role of transfusion**

Blood transfusion may restore the oxygen-carrying capacity in patients with SS disease with symptomatically lowered hemoglobin levels and may also serve to dilute circulating HbS-containing cells reducing the tendency to vaso-occlusion. Deviations from the patient’s steady-state values usually reflect a specific pathology requiring diagnosis, since this will determine the role of blood transfusion.

Symptoms of anemia are usually determined by the rate of fall in hemoglobin level which is most rapid in acute splenic sequestration (ASS; falls of 4–6 g/dL in hours), less precipitate the aplastic crisis (falls of 1 g/dL/day), and much slower in chronic renal failure and iron or folate deficiency. In chronic renal failure, the lack of erythropoietic drive to the bone marrow is followed by a gradual decline in hemoglobin which may respond to exogeneous erythropoietin (EPO), but is more readily treated by top-up transfusions to maintain oxygen carriage and relieve symptoms. In chronic hypersplenism, the low hemoglobin and the metabolic demands of the greatly expanded bone marrow may be relieved by chronic transfusion programs or more readily by splenectomy. Infections may suppress the bone marrow response, and infestations such as malaria may exacerbate hemolysis, so transfusions may have a supporting role while the underlying pathology is treated. Nutritional deficiencies of iron, folate, or possibly vitamin B₁₂ require treatment which is often followed by a prompt hematological response, although short-term transfusion may be necessary.

**Lowered hemoglobin levels – “top-up” transfusion**

**Acute splenic sequestration**

In the past, acute enlargement of the spleen associated with low hemoglobin and increased reticulocytes was a common cause of early mortality. In the Jamaican Cohort Study, attacks occurred as early as 3 months, were most common in the second 6 months of life with a cumulative probability rising from 0.255 by 2 years to 0.297 by 5 years, and were uncommon after 6 years of age. Events are more common...
in patients with low HbF levels,9 but the precipitating factors are unknown although events may be concurrent in twins.8 Recurrence is common, and after 2 attacks, further attacks may occur at shorter intervals, leading to a policy in Jamaica of prophylactic splenectomy after 2 attacks. In the absence of a clear etiology, prevention may not be possible, but mortality can be significantly reduced by parental education on early detection,8 which will allow time for transfusion during the acute episode. A single unit of blood, proportionately less for young children, is adequate, since this is frequently associated with a decrease in splenic size and release of sequestered blood which may raise the hemoglobin to unexpectedly high levels.

The use of chronic transfusion to prevent recurrence is controversial. Prophylactic chronic transfusion may have the advantage of restoring or improving splenic function,10 but it is unclear how best to monitor or when to stop such chronic transfusion. Well-controlled programs maintaining HbS levels below 20% may fail to prevent recurrences of ASS,11 and stopping transfusion at age 5–6 years when ASS becomes rare in untransfused patients may be followed by recurrent attacks.12 Considering these problems and the difficulties inherent in chronic transfusion, prophylactic splenectomy after 2 attacks of ASS appears to carry less risk, although the 2 approaches have not been assessed in controlled trials.13

Aplastic crisis

Defined clinically as a hemoglobin markedly below steady-state levels, usually to 2–5 g/dL, and absence of reticulocytes, the aplastic crisis almost always occurs due to parvovirus B19 infection.14 This organism destroys the early precursors in the bone marrow for 8–10 days exceeding the mean red cell survival in some patients, and hemoglobin levels fall by 1 g/day, leading to avoidable morbidity and sometimes death. In most cases, the outcome is benign and predictable, and transfusion may be performed as outpatients with follow-up for 3–5 days later to ensure that the expected reticulocytosis of the recovery phase has occurred. This is manifest by a rapid daily increase in reticulocyte count, which often reaches 30–40%, and the hemoglobin level is quickly restored to steady-state levels. Transfusion of a single unit of blood is adequate to maintain oxygen delivery during this period, but occasionally, the reticulocyte response is delayed and a second transfusion may be necessary. Human parvovirus is highly infective; 60% of SS patients seroconvert by 15 years of age,15 and SS siblings of an aplastic patient should be closely observed as approximately 80% of nonimmune siblings with SS disease are affected either simultaneously or within 3 weeks. Immunity to parvovirus appears to be lifelong, and recurrent attacks of B19 infection have not been described in otherwise uncomplicated SS disease.

**Chronic hypersplenism**

Sustained splenic enlargement with hemoglobin levels below and reticulocyte counts above steady-state levels may complicate SS disease, most commonly between the ages of 5 and 15 years, and occurred in approximately 5% of the Jamaican Cohort Study. Although the etiology is unknown, red cell survival in hypersplenism is further shortened from the usual range of 10–12 days to 2–3 days with hemoglobin levels of 3–5 g/dL, and reticulocyte counts generally exceeding 15%. Marked bone marrow expansion increases the metabolic demands, which compete with those of growth and may impair height velocity. The natural history of chronic hypersplenism in SS disease is poorly documented but may resolve spontaneously, although most patients require transfusion to maintain oxygen carriage and relieve symptoms while awaiting signs of spontaneous regression. Such cases are monitored in Jamaica with monthly assessments of spleen size, hemoglobin and reticulocytes, and height and weight with transfusion to relieve symptoms of diminished exercise tolerance. If there is no evidence of spontaneous regression within 6 months, therapeutic splenectomy is usually performed. Elective splenectomy, which avoids the cost and complications of chronic transfusion, has been universally beneficial in Jamaican experience. With pneumococcal prophylaxis which is given for at least 3 years, splenectomized patients do not appear to be more prone to overwhelming blood infection.16

**Chronic renal impairment**

A progressive glomerular fibrosis causes fall in glomerular filtration rates and lowering of EPO levels, and commonly contributes to mortality in patients with African forms of SS disease over the age of 40 years. Reticulocyte counts and hemoglobin levels fall slowly and are surprisingly well tolerated, with some patients not seeking medical attention until hemoglobin levels reach 2–4 g/dL. Patients may respond to exogeneous EPO, but treatment is expensive, difficult to administer, and because of the expanded bone marrow, unpredictable in its effects. The most economic therapy is regular top-up transfusion to relieve symptoms, which may be achieved with hemoglobin levels in the range of 6–8 g/dL. In Jamaica, many patients have been maintained for years with simple top-up transfusions, the frequency of which was determined by patient symptoms.
Transfusion to dilute the circulating sickle red blood cells

Transfusion may be beneficial in the prevention of stroke and in management of the acute chest syndrome, but there is no convincing evidence of benefit for therapy of the bone pain crisis, perioperative management, pregnancy, ASS, priapism, and chronic leg ulceration.

Central nervous system

A major pathology in childhood, stroke, affected approximately 10% of Jamaican Cohort by the age of 14 years, with a median age at first stroke of 6 years. The pathology is usually ischemic consequent on blockage of major cerebral, carotid, or vertebral vessels. Strokes recur in 50–70% of patients within 3 years, and chronic transfusion significantly reduces recurrent stroke compared to a non-transfused group. Transcranial Doppler (TCD) may detect stenosis of major blood vessels before strokes occur, and a randomized trial of chronic transfusion in patients with TCD evidence of cerebral stenosis showed a highly significant reduction of stroke, and of persistence of normal TCD indices, in the transfusion-treated group. Silent cerebral infarcts detectable by magnetic resonance imaging or “soft” neurological signs may also be predictive of further silent infarcts or strokes, and a randomized trial of transfusion over a median period of 3 years showed a marginal statistical improvement.

Although successful in the short term, chronic transfusion therapy raises many problems in long-term management such as its duration, the effects of stopping therapy, and lack of an optimal method of monitoring the effectiveness of chronic transfusion. Most programs are monitored by maintenance of HbS levels below 30%, but recurrences have occurred with HbS levels as low as 17–18%, and recurrent events, either overt strokes or fresh silent infarcts, occurred in 18 of 40 (45%) children on well-monitored transfusion programs, and stroke occurred after 11 years on transfusion, with HbS levels below 20%. The required duration of transfusion is also unclear, with recurrences occurring in 70% within 1 year of stopping transfusion programs of 1–2 years and in 50% within 1 year of stopping transfusion programs of 5–12 years, a higher frequency than suggested by the natural history data, suggesting that stopping transfusion actually raises the risk of recurrent stroke. The STOP 2 randomized study, which sought to assess the safety of stopping chronic transfusion after a minimal duration of 24 transfusions, was terminated prematurely because of the higher rate of high-risk TCD or occurrence of stroke in the transfusion-halted group.

These observations have led to suggestions that transfusion should be continued for life, raising the risks of long-term complications such as iron overload and red cell alloimmunization as well as with an estimated cost, computed in the year 2000, at US$40,000 per patient per year.

Acute chest syndrome

The acute chest syndrome has a complex pathology with elements of infection, infarction, fat embolism, and pulmonary sequestration and is a major contributor to deaths after the age of 2 years. Rapidly deteriorating dyspnea, clinical and radiological signs, and pulse oximetry characterize acute pulmonary sequestration which is a life-threatening emergency. Acute dyspnea associated with rapidly increasing, widespread pulmonary opacity may be dramatically reversed by exchange transfusion, which has been recommended within 48 hours. The incidence of acute chest syndrome and bone pain crisis was significantly reduced in transfused patients in the Stroke Prevention Trial.

Bone pain crises

Bone pain crisis, a characteristic feature of sickle cell disease, usually attributable to avascular necrosis of the bone marrow, is often precipitated by skin cooling, stress, and infection and is more common in the last 3 months of pregnancy and the immediate postpartum period. Bone pain crises were reduced in a prospective randomized trial of chronic transfusion in pregnancy, in the STOP trial, and in an earlier trial with less aggressive transfusion maintaining HbS levels below 50%. Although these observations are consistent with a beneficial effect of transfusion in the bone pain crisis, no data are available from a controlled clinical trial.

Perioperative management

The risks of serious morbidity and occasionally mortality which complicated surgery and anesthesia in the past have improved greatly, but it is unclear whether this is due to better general management or the increasing use of transfusion. Acute transfusion, repeated transfusion, and partial exchange transfusion have been advocated 10–15 days prior to or immediately before surgery, but comparison of transfusion techniques failed to reveal perioperative differences.

The first controlled trial found no perioperative differences between aggressive and conservative transfusion, but their protocol assumed that transfusion was essential and there was no non-transfusion arm in their study. The fundamental question of whether transfusion was necessary remained unanswered, since the morbidity was no greater in centers
Phylactic transfusion. There were no differences in fetal improvement that has occurred in centers not using prophylactic transfusion, but a higher level of blood exposure in chronic transfusion programs. These problems include iron overload, delayed hemolytic transfusion reactions, red cell alloimmunization, transfusion-acquired infections, maintaining venous access, and other side effects.

Iron overload

Each 300 mL unit of blood used in transfusion is estimated to contain 225 mg of iron. With the widespread use and greater duration of chronic transfusion, there is increasing concern on iron deposition in the liver and conduction bundles of the heart. Attempts to address this problem have included reducing the rate of iron accumulation by modifying transfusion methods and the use of chelation to remove iron present already in the body. Iron accumulation may be reduced by less rigorous target HbS levels, automated red cell exchange (erythrocytapheresis), which may delay or even reverse iron accumulation, and partial manual exchange transfusion.

Chelation is usually commenced when serum ferritin levels exceed 1000 ng/mL and traditionally depended on subcutaneous deferoxamine (DFO), which has a half-life in the plasma measured in minutes and has to be given by continuous subcutaneous injection overnight for 5–7 nights weekly. It is expensive, difficult to administer, and limited by compliance especially during adolescence. It was hoped that some of these disadvantages would be overcome by the oral chelator, deferiprone (DFP), with a 2-hour half-life requiring dosing 3 times daily and deferasirox (DFX; Exjade, Novartis International AG, Basel, Switzerland) which lasts 24 hours and requires daily administration. Both molecules are uncharged and theoretically would have easier access to intracellular iron compared to the charged molecule of DFO–iron complex. The excretion of DFP–iron complexes is predominantly through the urine, whereas the DFX is excreted through the stools. The features and side effects of the 2 oral chelators and DFO have been compared with those of an “ideal chelator” by Neufeld. The short-term safety of DFX appears acceptable, but data on longer term exposure are limited, although these concerns may have been allayed by a recent report of a 4-year follow-up. Despite the potential advantages of DFX, compliance may still be a problem because of the unpleasant taste. The management of transfusional iron overload in sickle cell disease has been recently reviewed.

Problems with transfusions

Occasionally, problems may occur with simple top-up transfusions in ASS or the aplastic crisis with acutely lowered hemoglobin levels, but these are greatly magnified with the higher levels of blood exposure in chronic transfusion programs. These problems include iron overload, delayed hemolytic transfusion reactions, red cell alloimmunization, and other side effects.
of the UK experience found major morbidity in 10 patients and death in 1 patient with sickle cell disease, and a retrospective analysis of 99 episodes in France reported death in 6%.

### Red cell alloimmunization

Alloimmunization is a major problem in chronic transfusion therapy in sickle cell disease. The overall rate in the Cooperative Study in the US was 18.6% and resulted from multiple antibodies in over 50% of alloimmunized subjects. Most antibodies were of the Rh (especially C, E), Kell (K), Duffy (Fy), and Kidd (Jk) groups tending to reflect differences between the predominantly Caucasian donors and Black recipients. Use of selected donors may reduce this problem, and closer racial matching of donor and recipient populations may have contributed to the lower alloimmunization rates of 12.9% reported from Brazil, and the 2.6% reported from Jamaica, although high rates of alloimmunization persisted in the US despite racial matching of donors and recipients. Alloimmunization rates have fallen with more extensive phenotype matching, rates of 18–76% being reported with standard matching by ABO and Rh groups, 5–14% with some phenotype matching for C, E, and K antigens, and 7% with more extended matching, and extensive matching reported no alloimmunization within 40 patients monitored at a single institution. To reduce alloimmunization, red cell phenotyping has been proposed prior to the first transfusion followed by matching at least the Rh and Kell groups, but a retrospective study found similar alloimmunization rates in US centers with or without extended phenotype matching. Even with extended phenotyping, alloimmunization continues, and the complex field of alloimmunization in sickle cell disease has been extensively reviewed. Further concerns on the significance of alloimmunization resulted from a recent study showing that alloimmunized patients were more prone to chronic pain, end-organ damage, and shortened survival. Post-transfusional monitoring may be helpful in the detection of new antibodies.

### Transfusion-acquired infections

These are inversely related to the diligence of diagnostic screening, and the risks of transfusion-acquired hepatitis B and C and of HIV and human T-lymphotropic virus have been greatly reduced, although the infections may still occur. Potential threats also persist with viral agents such as parvovirus B19. Recent review of transfusion safety in the UK indicated that the rates of infections had fallen substantially and that the greatest risk was clerical error. Malaria may be acquired by transfusion of malarial parasites, with a Ghanaian study finding parasites in 11% of transfused blood units, and cases may occur in traditionally non-malarial areas.

### Venous access

Chronic transfusion requiring venous access every 3–4 weeks clearly compromises the available veins, yet limited data are available on the frequency of implantable ports or catheters. These procedures accounted for 86 of 1033 (8.6%) of all surgeries in a trial of preoperative transfusion, and it is well recognized that these ports are prone to infection, thrombosis, and embolism, which may require their removal. A summary of earlier studies and a retrospective, single-institution study still did not provide prevalence data but reported that infections and thrombotic problems were more common among adults than children.

### Transfusion reactions

Non-hemolytic transfusion reactions used to occur with approximately 1% of blood units and in 10–40% of patients with repeat transfusions. Eighty-five percent of patients receiving more than 50 transfusions demonstrated antibodies to HLA or platelet antigens, but this has been markedly reduced with leukocyte-depleted blood.

### Supplies of suitable blood

In countries with well-developed blood bank services, suitable blood is generally readily available, although alloimmunization may limit the availability of compatible units. In countries with less well-developed facilities, obstacles to the provision of chronic transfusion programs include the limited availability of blood and lack of resources for extended cross matching. The latter may increase the risks of alloimmunization, although these may be reduced by the better racial matching of donors and recipients.

### Methods of transfusion

#### Simple transfusion

This is the method of choice for the acutely lowered hemoglobin levels in ASS and the aplastic crisis, and the chronically lowered hemoglobin levels in chronic renal failure. Packed red cells are usually used, and washed red cells or special filters may reduce transfusion reactions to white cells or platelet antigens.

#### Chronic transfusion program

Used increasingly for a variety of cerebrovascular complications, this requires transfusion every 3–4 weeks and is designed to maintain HbS levels below 30%. It may be
performed by recurrent simple transfusions, but attempts to reduce iron accumulation have led to exchange transfusions, which may be performed manually or by automated methods. Donor blood should be confined to the normal AA genotype, since although theoretically AS-genotype blood may be adequate in improving flow, it complicates the assessment of HbS levels.

**Manual methods**

A simple early method consisted of removal of 500 mL blood followed by infusion of 2 units of donor cells which achieved a 30% exchange in 90 minutes. More recent techniques generally use 2 intravenous lines with simultaneous or sequential withdrawal of SS blood and its replacement by donor AA cells. However, although manual procedures are effective in conducting exchange, they are tedious and time-consuming.

**Automated methods**

Machines performing centrifugal separation of red blood cells provide a rapid and efficient method of performing partial exchange transfusion. Two types of machines are in common use, the discontinuous- and continuous-flow models.

Discontinuous-flow cell separators act as batch processors, pumping blood from the patient’s arm, centrifuging it to separate red cells from the plasma, discarding the red cells, and returning the plasma to the patient along with donor cells. A standard procedure of 5–6 cycles achieves a 50% exchange in approximately 140 minutes. Continuous-flow separators use 2 intravenous lines, one extracting blood and the other returning the processed blood to the patient. Controlling rates of inflow and outflow ensures that the patient’s blood volume does not vary during the procedure. A 6-unit exchange can be completed in 70 minutes, a 72% exchange within 2.5 hours, and a 90% exchange in 3.5 hours. Such rapid exchange invariably results in a sharp increase in oxygen affinity which may cause problems, although it was well tolerated in other series. In summary, discontinuous-flow separators are small, mobile, simple to operate, relatively cheap, and require only 1 intravenous line. Continuous-flow separators are more complex and expensive, and more efficient but require 2 intravenous lines with large needles or catheters capable of carrying the large flows. Supplies for both cell separator techniques are considerably more expensive than manual methods but require less professional time.

**Acute exchange transfusion**

With rapidly progressive, life-threatening pathology, it may be necessary to conduct immediate exchange transfusion, and this is most commonly indicated in acute pulmonary sequestration or the acute phase of a stroke. Replacement of the HbS-containing cells in these situations may prevent the progression of pathology, and sometimes, dramatic reversals occur in clinical and radiological features indicating that some cases of acute pulmonary sequestration are a reversible vascular phenomenon.

**Choice of blood**

Chronic transfusion programs with the objective of lowering HbS levels may be more effective and simpler to monitor if donor blood is restricted to the HbAA genotype. Fresh blood should persist for longer in the recipient’s circulation and may reduce transfusion requirements. Concentrated young red cells (neocytes) have been used for transfusion programs in thalassemia but require large resources of blood from which to harvest the neocytes. The use of neocytes and a supertransfusion regimen has been shown to reduce iron accumulation in thalassemia, but no data are available on this approach in sickle cell disease.

**Optimal hemoglobin level**

Generally, patients with SS disease are well adapted to their steady-state hemoglobin level of 6–9 g/dL. In acutely lowered and symptomatic anemia, transfusion support should be the minimal required to maintain oxygen delivery. In the aplastic crisis due to parvovirus B19 infection, bone marrow activity is usually about to recommence by the time of clinical presentation, and in the author’s experience, a single unit is adequate. Special care is required in transfusion support for ASS when release of sequestered red cells from the spleen added to those given by transfusion may lead to unexpectedly high hemoglobin levels and sometimes resulting in bone pain. Chronic transfusion programs are usually monitored by the HbS level, and the total hemoglobin is a secondary consideration. Sudden changes in hematocrit may have serious implications for renal function, and excessive transfusion has caused death from cerebral hemorrhage.

**Conclusion**

Reducing the levels of HbS lessens the likelihood of pathology in sickle cell disease, and theoretically, regular transfusions of HbAA blood from birth would prevent the manifestations of the disease. In practice, the many problems with chronic transfusion make this unrealistic, and a balance must be struck between the advantages and disadvantages of chronic transfusion. The final position will be determined by many factors such as the ready availability of blood,
resources for extended blood group matching, facilities for performing exchange transfusion, maintaining of venous access, the availability of effective chelation, and the option of alternative therapies. Chronic transfusion programs are more readily achieved in major centers in the US and Europe but are a daunting prospect for developing societies because of the limited medical infrastructure and the large numbers of patients with sickle cell disease.

Indications for blood transfusion in the management of sickle cell disease present a spectrum from widespread acceptance, areas considered controversial, and those where there is almost no supporting evidence. Clearly beneficial is the use of top-up transfusion for acutely or chronically lowered hemoglobin levels, and convincing data support the use of chronic transfusion programs in the prevention of primary or secondary stroke. Controversial areas include the use of transfusion preoperatively, in the management of pregnancy, and in clinical complications such as chronic leg ulcers or priapism. Most controversial is the use or “top-up” transfusions at steady-state hemoglobin levels, some physicians believing that this may be beneficial. Inevitably, the clinically acceptable point in this spectrum is influenced by the sophistication of transfusion services and ready availability of blood. Despite convincing evidence, therapies such as chronic transfusion for the prevention of primary or secondary stroke may not be feasible in some societies because of the limited availability of blood and of resources available for iron chelation. It must be accepted that transfusion is not universally benign but carries with it risks that are multiplied with the transfusion loads associated with chronic transfusion programs. Whether these risks are justified must be determined by the natural history of the underlying complication and by the availability of alternative therapies. In communities where strokes are a major manifestation of sickle cell disease but the available resources make chronic transfusion unrealistic, hydroxyurea may offer an alternative therapy, and may also reduce transfusion requirements. In the prophylaxis of recurrent ASS or in the treatment of chronic hypersplenism, splenectomy offers a viable alternative to the dangers of chronic transfusion. It is true that these risks may be reduced by extended phenotype matching, at some cost, but even then, is it necessary to take these risks if alternative therapies are available? More information is needed on the natural history of the complications of sickle cell disease and of alternative therapies before the role and risks of transfusion therapy can be clearly defined.

The African forms of sickle cell disease have more severe features than those with the Asian haplotype where frequent alpha thalassemia and persistence of HbF may protect against some of the serious early pathologies of the disease. More information is needed on the Asian haplotype, before the role of transfusion can be determined in that population, and transfusion in the steady state requires justification. As the Asian haplotype of SS disease becomes better documented and understood, the role of transfusion therapy in its management will be more readily defined.

Disclosure

The author reports no conflicts of interest in this work.

References


