Erythrocytapheresis: Do Not Forget a Useful Therapy!

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Key Words
Erythrocytapheresis · Red blood cell exchange · Sickle cell disease · Exchange transfusion

Summary
In patients with pathologically altered erythrocytes, red blood cell exchange is a very efficient therapeutic measure without important side effects. With increasing migration and relocations, more patients with e.g. severe malaria or sickle cell anemia have to be treated. In minor or bidirectional ABO-mismatched stem cell transplantations after reduced intensity conditioning, hemolysis can be prevented by prophylactic erythrocytapheresis. Other rare indications for red blood cell exchange are advanced erythropoietic protoporphyria and babesiosis. Sickle cell anemia can be treated with hydroxyurea. Transfusions are administered when necessary, but this results in iron overload in the long term. An expensive but safe and very efficient treatment alternative is red blood cell exchange. In cases with stroke, acute chest syndrome and other severe complications, erythrocytapheresis reproducibly breaks the vicious circle of sickling and increasing oxygen deficiency. At the same time one can aim at an exact end hematocrit. In severe malaria, erythrocytapheresis both reduces parasite load to the designated extent and constitutes reduced oxygen transport capacity without serious adverse effects. Here we describe our experience of erythrocytapheresis in long-term prophylaxis of complications in sickle cell anemia and sickle cell thalassemia patients. The documentation of improved iron balance was carried out by liver susceptometry.

Schlüsselwörter
Erythrozytapherese · Austauschtransfusion · Sichelzellanämie · Erythrozytenaustausch

Zusammenfassung
Introduction

Erythrocytapheresis, also referred to as red blood cell exchange, is a procedure rarely thought of. The use of modern cell separators, the introduction of leukodepletion in blood products, and the increasing safety of red blood cell concentrates have made this treatment modality very effective, comfortable and safe.

The increasing number of persons with African, Asian and Mediterranean origin in our country has led to an increased incidence of sickle cell anemia and sickle cell thalassemia patients. The rising number of trips to malaria regions obliges to treat malaria regularly.

Repetitive transfusions cause iron overload. This has so far been treated by subcutaneous application of chelators with very poor compliance. The advent of oral chelators mitigates this problem. However, there still are indications where the increased blood viscosity and insufficient relative reduction of pathologically altered red blood cells needs better and faster treatment [1–3].

Sickle cell anemia is caused by homozygous gluiamond (hemo-
globin S; HbS) mutation. The disorder is recessively inherita-
able, heterozygotes for gluiamond and gluodynamics (hemo-
globin SC) suffer from sickle cell disease, too. Similar clinical problems are found in patients with sickle-beta-thalassemia. The course of these defects is highly variable, and progress in predicting complications is under progress.

Children who have suffered from one ischemic cerebrovascu-
lar accident have a very high risk of relapse that can be signifi-
cantly lowered by transfusions [4–12]. But transfusion-in-
duced increase in blood viscosity can aggravate perfusion problems. Erythrocytapheresis is a very safe and efficient way of both reducing blood viscosity and improving oxygen delivery to the tissue [1, 2].

Hemotherapy for sickle cell disease complications is divided into treatment of acute crises such as stroke and acute chest syndrome and prevention of complications and relapses. In crises hemoglobin polymerization causes sickling of red blood cells and leads to vasoocclusion and perfusion problems. Erythrocytapheresis is a very safe and effective technique of both reducing blood viscosity and improving oxygen delivery.

We performed erythrocytapheresis using the Cobe Spectra cell separator with the standard erythrocytapheresis program in the older patients with veno-venous access; in small chil-
dren blood was given back via port. At first an ACD-to-whole blood ratio of 1:14 was used, later we gave ACD at a rate of 1:17 with the maximally tolerated blood flow.

Exchange transfusion of about one patient’s blood volume increasing the hematocrit by maximally 0.02 in anemic patients.

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patient mild citrate reactions were noted. The first was overcome by catheter implantation and the latter by reduction of the citrate rate to 1:17 and calcium substitution into the return line. Both stroke patients recovered excellently and did not suffer from any relapse in spite of high risk. Iron overload was tolerable. The one patient with many transfusions before red cell aphereses had subcutaneous chelation therapy, but she abandoned that because of side effects. With the availability of oral chelators, this should be started again in spite of her compliance. The two patients with pain crises showed clinical complications every time they exceeded routine apheresis intervals because HbS rose too high.

**Results**

**Patient 1**
Patient 1 is a young woman born in 1983 in Africa. She has always been transfusion-dependent. At first treatment at our hospital in July 1990 she suffered from a hematocrit of 15%, hepatitis C as well as a lack of folic acid and vitamin B12. In June and November 1991 she had two pain crises. In August 1992 she suffered from vascular obliterations and received two red blood cell concentrates. In the following year a program of 6–12 weekly transfusions was instituted. In February 1995 epistaxis was observed, since September 1995 red cell exchanges were carried out every 6–12 weeks and continued until 2000. In 1996 subcutaneous Desferal was started to reduce iron overload. Two years after the start of chelation sudden hearing loss occurred, probably as a side effect of this treatment, and Desferal was stopped. In 2000 hydroxyurea was started and aphereses were stopped, but the patient suffered from further pain crises. In 2001 she received red cell transfusions during pregnancy. In February 2002 erythrocytaphereses were started again and continued until today. So far she has undergone 62 red blood cell aphereses. The patient's compliance is variable. In the course of the last year she suffered from two pain crises when intervals between aphereses became too long. Ferritin as an inflammatory protein rose and decreased again as the hepatitis became clinically more stable. Her ferritin level as a marker of inflammation was very high and extremely variable at periods with greater hepatitis activity (fig. 1). Liver susceptometry showed increased LIC, but less than would be expected with blood transfusions instead of red cell exchanges (fig. 2). She always has signs of hemolysis and inflammation such as elevated reticulocytes, reduced haptoglobin, and high LDH and leukocyte levels. In spite of these symptoms, iron overload and deterioration of complications such as pain crises and osteonecrosis could be delayed. Iron overload is clearly less problematic than in simple blood transfusion. Even hemolysis of sickle cells was mitigated.

**Patient 2**
Patient 2, a young man born in 1987, suffers from combined sickle cell-beta thalassemia and has been symptomatic since his 4th year of life. Until the age of 9 years he had received 4 red blood cell concentrates. In February 1998, at an age of 10 years, he presented in our hospital with a pain crisis and received 2 red blood cell concentrates. In April of the same year automated red blood cell exchanges were started and continued until now at 4-weekly intervals. The patient's compliance has decreased during the course of adolescence, and every time the intervals between blood exchanges became too long the patient suffered from pain crises that lasted for several days, improving only slowly after erythrocytapheresis. Overall he was treated with 82 red blood cell aphereses so far. Levels of ferritin, an acute phase protein, first rose, but decreased during adolescence, perhaps reflecting emptying iron stores due to growth. LIC was not as high as in comparable patients with transfusions only, and the number of complications could be kept low by repeating aphereses at short intervals. LIC was...
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Fig. 2. a LIC by SQUID biosusceptometry under continuous erythrocytapheresis treatment in 4 patients with sickle cell disease. Median LIC for SCD under blood transfusion treatment according to [15]. b Ferritin-to-LIC ratio under continuous erythrocytapheresis treatment in 4 patients with sickle cell disease. Expected interquartile range for SCD under blood transfusion treatment according to [15].

Fig. 3. HbS before and after erythrocytapheresis treatment in 4 patients with sickle cell disease.
highest in 2000, now it has become normal with elevated ferritin levels. LIC and ferritin levels are shown in figures 1 and 2. The HbS levels have been kept in a tolerable range as shown in figure 3. In 2004 MRT was carried out because of continuing pains in the legs. Traces of deep vein thromboses and aseptic bone necroses in both acetabuli and femora were seen. Clearly, without apheresis this patient would have many more clinical complications and iron overload due to the necessary transfusions.

**Patient 3**

Patient 3 was born in December 1998. Sickle cell disease was diagnosed in 2001. He suffered from stroke in January 2003 ensued by left-sided hemiparesis and convulsive versive seizures that had to be treated with carbamazepin. Hemosiderosis at that time was quantified by liver susceptometry (fig. 2). The left leg was operated on several times to alleviate spastic paresis consequences. Meanwhile, the patient has learned to walk again in spite of his overweight (54 kg, 141 cm). Four months after the stroke a 4- to 6-weekly red cell exchange program was started in May 2003 as secondary stroke prophylaxis. Due to poor venous access a port-a-catheter was implanted into the left jugular vein in August of the same year. Up to now 46 erythrocytaphereses were carried out every 4-6 weeks without side effects. Hemosiderosis did not progress. Cerebral blood flow as assessed by transcranial Doppler sonography was normal in posterior cerebral arteries; the other arteries were not evaluable. Residues of the stroke are seen in the EEG. LIC has increased from 432 mg/g liver in March 2005 to 910 mg/g liver in November 2006, ferritin is 561 mg/l. Overall the iron load is tolerable as shown in figures 1 and 2. No erythrocytapheresis complications were seen in this high-risk patient, and even iron chelation therapy has not become necessary.

**Patient 4**

Patient 4 was born in February 1999, and homozygous sickle cell anemia was diagnosed in June 2002 during pneumococcal sepsis. In April 2002 he suffered from a stroke and recovered quite well. Transcranial Doppler sonography was normal in posterior cerebral arteries; the other arteries were not evaluable. Residues of the stroke are seen in the left fronto-parietal region. Since July 2003 he received regular red blood cell aphereses every 4–5 weeks. Ferritin values varied highly and are now in a tolerable range (fig. 1). LIC is still in the range where no chelation therapy is necessary. No clinically relevant problems occurred. All 46 erythrocytaphereses carried out so far were tolerated well. Since 2004 he complained of abdominal pain and flatulence at the end of each apheresis procedure. As a consequence we reduced the citrate ratio from 1:13 to 1:17 and substituted calcium during every treatment. After that no more side effects have been observed. At 9 years of age he is 136 cm tall, with a weight of 56 kg and a head circumference of 57.8 cm.

**Discussion**

Problems in sickle cell disease arise from deficiencies in oxygen delivery to the tissue due to erythrocyte deformation, hemolysis, and inflammation. Increase in blood viscosity leads to perfusion problems, mainly in large vessels. Hemolysis adds to this problem by scavenging of the vasodilator NO released by endothelial cells. Inflammation and increased cell adhesion of erythrocytes as well as leukocytes and platelets contribute to vasoocclusion, especially in small vessels, during crises. Blood viscosity of oxygenated sickle cell blood is 1.5-fold that of normal blood, but increases to 10-fold upon deoxygenation. This increase is the higher the faster the deoxygenation occurs. High hematocrit contributes to increasing viscosity. Transfusion alone further increases blood viscosity and limits relative reduction of HbS.

Blood usually is transfused at a speed of about 2 h per red cell concentrate. Elevating hemoglobin above 10 mg/dl often causes a too steep increase in blood viscosity [3]. Moreover, red cell concentrates are usually transfused slowly, and a standard transfusion of two units takes at least 3 h. This limitation is easily overcome by apheresis, up to 10 red blood cell concentrates can easily be exchanged in less than 3 h without relevant side effects, thus giving the possibility of immediate improvement in case of crisis and providing a more comfortable procedure in patients treated prophylactically. Erythrocytapheresis is the only way to reduce quickly and in a comfortable way for the patient the concentration of HbS-containing red cells. As shown in figure 3, in our patients an immediate reduction of HbS below 30% as postulated by Pegelow et al. [12] could be achieved by erythrocytapheresis. The major advantages of erythrocytapheresis are the decrease in blood viscosity in parallel to the increase in oxygen transport capacity. At the same time defective red blood cells are replaced; thus hemolysis and ensuing inflammation, cell adhesion and NO antagonism are reduced [16].

During our prophylactic erythrocytaphereses we aimed at keeping the hematocrit rather stable and tried not to increase hemoglobin values much above the levels the patients were adapted to, usually ranging from 0.30 and 0.35. This relatively low hematocrit contributes to prevent risks related to high viscosity. Actually, our targets for prophylactic erythrocytapheresis are similar to those expected in case of stroke, acute chest syndrome and acute multiorgan damage syndrome, where a decrease of hematocrit to 0.3 and an immediate reduction of HbS red blood cells should be aspired. Despite the use of a great number of red cell concentrates, a long-term erythrocytapheresis program is extremely useful to minimize iron overload although it has little or no effect on a pre-existing iron overload related to a conventional transfusion program. At the same time free hemoglobin and NO scavenging are reduced [16], restoring normal vasodilatation and endothelial function.
Our secondary prophylaxis patients are all at highest risk to develop severe sickle cell disease complications. Since they are treated by erythrocytaphereses, they have not suffered from severe or irreversible problems. The young stroke patients have made fair progress in improving their stroke complications; both have started to grow normally, which can often be a problem in this patient group with low hemoglobin levels [17]. Transcranial Doppler blood flow velocities in the evaluable arteries are normal by now in both boys. The two patients with pain crises regularly suffer from relapse when intervals between treatment sessions were extended too much and improve within days after red blood cell apheresis. Biopspectrometry shows in all patients an iron overload that is much lower than in transfused patients. They are all adequately treated with a good quality of life and do not need chelators even after such a long disease duration.

Another indication for prophylactic red blood cell exchange is the prevention of severe immune hemolysis in minor and bidirectionally ABO-mismatched alloimmune peripheral blood progenitor cell transplantation after reduced intensity conditioning. Incidence and severity of this life-threatening complication can be reduced by prophylactic apheresis [18]. Erythrocytapheresis is helpful for the treatment of various crises. It leads to immediate improvement of symptoms in severe malaria and babesiosis [19, 20] and has also been successfully carried out for the prevention of alloimmunization against D+ antigen in combination with intravenous immunoglobulin [21]. Autoimmune hemolytic anemia is caused by circulating autoantibodies against red blood cells. Acute symptoms can be relieved by exchanging incompatible autologous red blood cells against compatible donor erythrocytes [22].

In erythropoietic protoporphyria ferrochelatase, the final enzyme in heme formation, is lacking. So protoporphyrin is deposited in the tissue, and blood protoporphyrin levels are elevated. Plasma exchange can reduce protoporphyrin levels, but red blood cell exchange has been reported to be more efficient because its concentration is severalfold higher inside red blood cells [23]. Preoperative red cell exchange may also be useful in patients with severe sickle cell disease to prevent complications from operation or anesthesia [24]. Nowadays, the decision to include a patient with sickle cell disease in an erythrocytapheresis program is based mainly on the previous occurrence of a major complication such as stroke, acute chest syndrome, or multi-organ damage syndrome. However, there is a great amount of recent works trying to identify risk factors of such complications in patients with sickle cell disease, e.g. intensity of hemolysis is correlated to leg ulcers [25]. Factor XIII, factor V Leiden and fibroblast growth factor 2 disturbances can all add to sickle cell complications.

Since 2004 various articles have related additional genetic risk factors and sickle cell complications: TNF-α promoter polymorphisms, IL-4 receptor and adrenergic β2 receptor are related to large vessel stroke, and VCAM1 and low density lipoprotein receptor polymorphisms show correlation to small vessel stroke [27]. α-Klotho, annexin-2 and bone morphogenic protein 6 are involved in vascular disease as well as bone calcium regulation and metabolism and bone morphogenesis and have also been shown to be risk factors for sickle cell disease complications [26–33]. Therefore, it is likely that in the future erythrocytapheresis programs will be more often initiated as a primary prophylactic intervention.

References


