CASE REPORT

Unusual splenic sinusoidal iron overload in sickle cell/haemoglobin D-Punjab disease

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Sickle cell/haemoglobin D-Punjab disease (SD disease) is a rare sickling disorder. The clinical phenotype closely resembles that of sickle cell anaemia. We describe the case of an 11 year old boy with SD disease on a regular blood transfusion programme. Splenectomy was performed at the age of 10 for progressive hypersplenism. Despite a history of regular iron chelation treatment, splenic histology showed evidence of profound iron overload. The dangers of iron overload are well recognised in children with thalassaemia, but less so in other hereditary haemolytic anaemias.

CASE HISTORY

Our patient is the second child of unrelated parents, both originating from southern Turkey. He had presented to his local hospital in Turkey at the age of 2 months with anaemia and hepatosplenomegaly. A diagnosis of ‘Mediterranean anaemia’ was made and a transfusion programme was begun. At the age of 2 years he moved to the UK with his family who brought him to our hospital requesting transfusion. At that time, he was clinically well with mild jaundice, splenomegaly (5 cm, palpable), and hepatomegaly (3 cm, palpable). Haemoglobin electrophoresis showed bands in the positions of haemoglobins S (24%) and A. Parental studies showed his father to be a carrier of haemoglobin S and his mother to be a carrier of haemoglobin D-Punjab. Subsequent studies, when he was not receiving transfusions, confirmed that the haemoglobin A was transfusion derived, and a diagnosis of SD disease was made. This was confirmed by molecular analysis, which showed haemoglobin D-Punjab (Los Angeles) (β121 glu→gln). Bone marrow examination at presentation showed sickle cells, erythroid hyperplasia, and mild dyserythropoiesis. Electron microscopy showed no evidence of congenital dyserythropoietic anaemia. There was no evidence of α or β thalassaemia trait or a red cell membrane defect in the patient or other family members. Glucose-6-phosphate dehydrogenase and pyruvate kinase assays were normal.

Transfusions continued and iron chelation with subcutaneous desferoxamine was started (escalating doses up to 60 mg/kg/day, given over 10–12 hours, six nights/week). Despite chelation treatment, the child’s serum ferritin rose to 3000 μg/litre. At 9 years of age, his transfusion requirement rose rapidly to 30 ml of packed cells/kg every three to four weeks. This was accompanied by increasing splenomegaly, mild leucopenia, and thrombocytopenia. Isotope imaging with technetium-99m labelled red blood cells showed an increased splenic red blood cell pool (19.9% of total red blood cell volume), with filling defects suggesting previous splenic infarction. Splenectomy was performed at the age of 10 years. He has remained transfusion independent since then, with a haemoglobin concentration of 70–80 g/litre. Moderate vasoocclusive crises developed on stopping the transfusion programme.

SPLENIC PATHOLOGY

The spleen weighed 405 g (mean normal spleen weight for a 10 year old boy, 110 g; SD, 30) and showed several areas of recent infarction. The cut surface had an unusual tan colour. Histologically, the white pulp appeared reactive but was otherwise unremarkable. The red pulp was expanded with widened cords and prominent sinuses. Sickle cells were visible in infarcted areas (fig 1A). Sinusoidal (but not capillary or larger vessel) endothelial cells showed striking haemosiderin loading, which was confirmed by Perls’ staining (fig 1B) and ultrastructural examination (fig 2). Cordal and intrasinusoidal macrophages showed moderate iron loading.

DISCUSSION

SD disease is a sickling disorder that is at least as severe as sickle cell anaemia. The usual clinical features of this condition were completely suppressed in our patient by regular transfusion from an early age.

Macrophage iron loading is well recognised in patients with haemolytic anaemia or transfusional iron overload. However, the sinusoidal endothelial iron loading in this patient is highly unusual. Possible explanations include direct phagocytosis of sickled cells or iron uptake via transferrin receptors.

In support of the first theory, cells from patients with sickle cell disease are known to undergo sickling in the hypoxic environment of the spleen, and sickled cells adhere to upregulated adhesion molecules on endothelial cells. Although phagocytosis in sickling disorders has not been demonstrated, splenic sinusoidal endothelial cells have been shown to phagocytose red blood cells in cases of hereditary spherocytosis and autoimmune haemolytic anaemia. Direct uptake of iron by endothelial cells is also possible, because...
transferrin receptors have been demonstrated on splenic sinusoidal cells.4 It has been speculated that splenic sinus lining cells (also known as littoral cells) have hybrid properties of endothelial cells and macrophages. In addition to transferrin receptors, splenic sinusoidal endothelial cells and littoral cell angiomas derived from them have been shown to express CD68.5 This has been taken as evidence of a hybrid endothelial–macrophage cell type. Nonetheless, splenic sinusoidal endothelial cells are undoubtedly highly unusual, apparently unique in the human body. Among their unusual properties as endothelial cells, they lack expression of CD34 but express CD8 and BCL-2. Intense, selective haemosiderin accumulation by these cells in our patient’s spleen is certainly compatible with phagocytic potential, although we found no visible evidence of ingested sickle cells within the sinusoidal endothelium.

Recent studies suggest that alternative treatments such as hydroxyurea are useful in ameliorating symptoms in SD disease and in sickle cell anaemia.* Although transfusion programmes abolish symptoms in these patients, this child’s case illustrates that the dangers of iron overload should always be remembered.

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