Clinical Reports

Partial splenectomy for massive splenomegaly secondary to Gaucher’s disease

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Summary: A 16 year old girl with Type 1 Gaucher’s disease presented with massive splenomegaly, hypersplenism and abdominal discomfort. Traditionally hypersplenism has been treated with splenectomy, but this results in a high incidence of overwhelming sepsis and accelerated sphingolipid deposition in both liver and bone. A 90% partial splenectomy was therefore performed leaving a fully vascularized inferior segment of the spleen and resecting 5.8 kg of splenic tissue. The patient made an uneventful recovery with a marked improvement in her haematological parameters and general condition.

Introduction

Gaucher’s disease is an autosomal recessive genetic disorder resulting in a deficiency of B-glucocerebrosidase which leads to an accumulation of glucocerebroside in the reticuloendothelial system and bone marrow. The principal clinical presentations of patients with Gaucher’s disease are hepatosplenomegaly and bone pain. Massive splenomegaly with hypersplenism is therefore a common presentation in Type 1 Gaucher’s disease (adult form) and in the past this has been treated by splenectomy. However, progressive hepatomegaly and accelerated bone disease have been reported after this procedure, as have overwhelming septic complications that can be fatal. These results have suggested that patients with massive splenomegaly secondary to Gaucher’s disease may benefit from a partial splenectomy. A partial splenectomy procedure was first described in 1869, and subsequently has been utilized following trauma, splenic cysts and haematological disorders. Partial splenectomy for Gaucher’s disease has been reported in Israel and the USA, but we now report the first documented case in the British literature in one of the largest spleens in which this procedure has been attempted.

Patient and methods

A 16 year old female weighing 35 kg presented with progressive symptoms of lethargy, abdominal pain and amenorrhoea and was found to have massive splenomegaly to the level of her pelvis (Figure 1). Four years previously she had been diagnosed as having aseptic femoral head necrosis. Peripheral blood results were haemoglobin 7.3 g/dl, white count $2.7 \times 10^9$/l and platelets $44 \times 10^9$/l. Bone marrow trephine biopsy showed extensive infiltration by Gaucher’s cells and the diagnosis of Type 1 Gaucher’s disease was confirmed by reduced plasma $B$-glucosidase activity and increased levels of acid phosphatase and angiotensin converting enzyme. X-rays of her skeleton showed signs of old avascular necrosis of the right femoral head (Figure 2) and abnormal trabecular pattern of both proximal femora.

One possible treatment option was allogeneic bone marrow transplantation but this could not be undertaken since her unaffected sister was HLA incompatible. Because of symptoms associated with her large spleen, it was decided to proceed to partial splenectomy to try and avoid rapid deterioration of her skeletal problems.

Pre-operatively the patient received pneumococcal vaccine, a transfusion of platelets and cefuroxime prophylaxis. Laparotomy revealed massive splenomegaly, no splenunculus and moderate hepatomegaly. The initial operative manoeuvre was to open the lesser sac and place a rubber sling around the splenic artery as it ran along the upper surface of the pancreas. The spleen was then mobilized dividing the spleno-renal ligament and omental and diaphragmatic attachments. The short gastric vessels were divided and the spleen delivered into the wound. Hilar dissection then started at the upper pole, ligating and dividing all vessels except for the most inferior artery and
Figure 1 Massive splenomegaly.

Figure 2 Right sided aseptic femoral head necrosis secondary to Gaucher’s disease.

Figure 3 The spared inferior artery and vein to the lower pole of the spleen being elevated by the artery clip, and the line of demarcation (arrowed).

vein supplying the lower pole of the spleen (Figure 3). The devascularized portion of the spleen became less tense and a clear line of demarcation was seen. The spleen was divided along the demarcation line and a 90% splenectomy performed resecting 5.8 kg of spleen (14% of body weight: normal splenic/body weight ratio 0.3%). Pressure was applied to the cut surface and the spleen was wrapped in a mesh of absorbable polyglycolic acid with a layer of the haemostatic substance spongian over the cut surface. The spleen was then sutured to the diaphragm in 3 sites using the mesh to prevent torsion.

The patient made an uneventful recovery with minimal drainage. She was discharged on the tenth day following surgery.

At follow-up the patient has made excellent progress. Her haemoglobin, platelets and white cell count 6 months later were 12.9 g/dl, 130 × 10⁹/l and 8.2 × 10⁹/l respectively and her weight 50 kg. Her periods are regular and normal. Clinical examination and CT scan give no indication of progressive enlargement of the remnant at 7 months and an isotope scan shows normal splenic perfusion.

Discussion

The management of patients with massive splenomegaly and hypersplenism secondary to Type I Gaucher’s disease has been unsatisfactory. In the past the only option was to perform a splenectomy, but this was often accompanied by a high incidence of sepsis and even death.⁴ The immune system is further embarrassed in these patients by the infiltration of the remaining reticuloendothelial system by the deposition of the Gaucher cells, placing these patients at even greater risk of sepsis following splenectomy than those undergoing such surgery for trauma, immune thrombocytopenic purpura or hereditary spherocytosis. Furthermore, after splenectomy in patients with Gaucher’s disease, there is often a progression in the hepatomegaly and in bone disease; in this patient, who was already having trouble with her right hip due to aseptic femoral
head necrosis, this would have been disastrous. Other treatment options were therefore considered.

Allogeneic bone marrow transplantation could not be undertaken since her unaffected sister was HLA incompatible. However, even if feasible, this would not have addressed the issue of her abdominal pain and difficulties in moving due to the splenic bulk. Arterial embolization can be considered but has to be done in stages, may be accompanied by a high fever and marked discomfort due to splenic infarction and does nothing initially for the bulk of the spleen.

Partial splenectomy in animals has been shown to inflict far less of an onslaught on the immune system than total splenectomy, and in Gaucher’s disease would decrease the bulk without increasing significantly the hepatomegaly or bone disease.

The actual procedure of partial splenectomy for such a vast spleen is a daunting prospect. However, with meticulous surgical technique, it is a feasible and safe procedure. In this patient no post-operative complications occurred, but in the cases reported from Israel and the USA a few complications have been reported including torsion of the splenic remnant, haemorrhage, necrosis of the remnant and left-sided pleural effusion.

Partial splenectomy in these patients undoubtedly improves the state of hypersplenism, and is also accompanied by an increase in body weight, muscle bulk and tone of abdominal musculature. It is of interest to note that our patient commenced her menses within 2 months of surgery and one similar case has also been previously reported in which a 20 year old female commenced having normal periods within 2 months of partial splenectomy, while in children it appears to result in an improvement in the growth curve.

The rate at which the remnant may enlarge is unknown and as yet this procedure has not been attended by a long enough follow-up adequately to assess the ultimate fate of the remaining spleen. Follow-up scans of up to 1 year have shown some slight increase in remnant size but nothing of clinical significance. It has been the objective of other workers to leave a remnant roughly the size of a normal spleen, but in this case the enormous size of the spleen (5.8 kg) dictated that the size of the remnant depended on the vascular supply. Thus the remnant was that part of the spleen that was adequately vascularized on only a single vessel. To have removed more would have run this risk of potential haemorrhagic complications.

This case therefore demonstrates that a partial splenectomy is a feasible possibility even with enormous spleens. The role of this procedure in Gaucher’s disease remains to be proven, but thus far looks most encouraging.

References

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doi: 10.1136/pgmj.67.794.1072

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