Massive gastrointestinal haemorrhage associated with ileal lymphoid hyperplasia in Gaucher’s disease

D.R. Jones, J. Hoffman, R. Downie and M. Haqqani

Walton Hospital, Liverpool, Merseyside, UK

Summary: A case report of adult type I Gaucher’s disease is described, with profound terminal ileal haemorrhage in association with ileal lymphoid hyperplasia. Peroperative ileal endoscopy localized the segment involved allowing an appropriate resection and successful outcome for the patient.

Introduction

Adult or type I Gaucher’s disease is characterized by hepatic and splenic accumulation of glucocerebroside, an intermediate product of membrane lipid metabolism. Bone involvement is frequent but only rarely are other organs affected; cerebral involvement is not a major feature and life expectancy is usually normal unlike the neuronopathic varieties, types II and III.

The pattern of disease is heterogeneous, probably due to variation in mutation in the gene for the affected enzyme: lysosomal glucosylceramide-beta-glucosidase.2

Blending complications arise as a result of thrombocytopenia associated with hypersplenism; the usual manifestations are petechial skin haemorrhage, bruising and epistaxis. Gastrointestinal haemorrhage is uncommon, haemopericardium has been reported3 as have oesophageal varices.4 We describe a case of massive gastrointestinal haemorrhage in association with ileal lymphoid hyperplasia in a patient with adult Gaucher’s disease.

Case report

A 47 year old male patient with adult, type I Gaucher’s disease was admitted following a profound rectal loss of blood whilst driving. He had had no previous significant rectal bleeding but was due to be admitted electively for excision of haemorrhoids. There had been no warning other than a sudden urge to defaecate.

At the age of 4 he had undergone splenectomy for splenomegaly and had been treated conservatively for bone deformity of the knees in childhood. A Billroth I gastrectomy was undertaken at the age of 22 and a Meckel’s diverticulum with an adhesive band had been excised some 17 years later. In the previous 9 years to his present admission he had a further 3 admissions for small bowel obstruction presumed to be adhesive in origin.

On admission he was pale, hypotensive, with a blood pressure of 80/40 mmHg and a tachycardia of 100/minute. Full blood count showed a haemoglobin of 9.3 g/dl and a normal platelet count of 234 × 10⁹/litre. Examination of the blood film showed the changes of hypoplasenism with nucleated red cells. Prothrombin time and kaolin-cephalin clotting time were not elevated. Serum alkaline phosphatase and transaminase enzymes were normal as were plasma electrolytes and urea.

Following resuscitation and cessation of bleeding, arrangements were made for an elective colonoscopy. On the day of the examination he had a further massive rectal loss of dark blood and underwent urgent sigmoidoscopy and colonoscopy. Other than haemorrhoids no identifiable mucosal lesion was visualized up to the level of the hepatic flexure at which blood was seen to be oozing from a more proximal site.

Immediate laparotomy revealed gross hepatomegaly but no other abdominal mass. There was blood within the lumen of the whole of the colon and 10 cm of the terminal ileum although the bowel was palpably normal. Peroperative flexible endoscopy with a gastroscope inserted through a small enterotomy in the ileum localized the site of bleeding to the terminal 20 cm of ileum up to the ileocaecal valve. The right colon and affected terminal ileum was resected to include any possible right colonic angiodysplasia. Postoperative recovery was uneventful and the patient was free of symptoms at review some 6 weeks later.

Correspondence: D.R. Jones, M.Ch., F.R.C.S., 556 Pensby Road, Thingwall, Wirral L61 7UE, UK. Present address: Vascular Studies Unit, Bristol Royal Infirmary, Bristol, UK. Accepted: 19 November 1990
Examination of the opened specimen showed the mucosal surface to be covered in a carpet of pale nodules approximately 1 mm diameter but more confluent nearer the ileocaecal valve. An area of telangiectasia near the ileocaecal valve marked the site of origin of haemorrhage. No mucosal abnormality was identified within the colon.

Histology of the ileal mucosa showed widespread reactive benign lymphoid hyperplasia with submucosal oedema and haemorrhage. Gaucher cells were not seen within the specimen. No vascular abnormality was apparent within the bowel wall (Figure 1).

Microscopy of a needle core liver biopsy showed no evidence of cirrhosis but engorgement of the parenchyma with typical Gaucher cells and distortion of the normal liver architecture.

Discussion

Haemorrhagic complications of Gaucher's disease are more usually associated with thrombocytopenia due to splenomegaly and bone marrow infiltration with glucocerebrosidase. Immune thrombocytopenia sufficient to cause retroperitoneal haematoma has been reported in one patient with Gaucher's disease and was suspected as the cause of thrombocytopenia in another. Gastrointestinal bleeding secondary to thrombocytopenia is uncommon although buccal mucosal petechiae have been noted. Patients may have low platelet counts for many years before bleeding diatheses develop. Moderate hepatic dysfunction secondary to accumulation of glycolipid causes elevation of blood liver enzymes and abnormal radioisotope uptake scans. The partial thromboplastin time may be prolonged due to an apparent deficiency of several clotting factors, particularly factor IX, in vitro. Hypofibrinogenaemia has not been reported. The case described above had a life-threatening haemorrhage some 40 years following splenectomy in the presence of a normal platelet count and normal clotting studies.

The liver is invariably involved in type 1 Gaucher's disease although development of cirrhosis is infrequent. Frank portal hypertension has been reported but is uncommon although oesophageal varices leading to gastrointestinal haemorrhage can occur and rarely may be massive and fatal.

Lymphoid hyperplasia of Peyer's patches does not feature in reports of Gaucher's disease although Gaucher cells are frequently found in Peyer's patches. Peyer's patch lymphoid hyperplasia is an uncommon finding but is associated with acute and chronic inflammatory conditions. It may give rise to ulceration and bleeding most notably in untreated typhoid infection.

The diagnosis and management of caecal or ileal haemorrhage presents problems, particularly when the bleeding is massive and life-threatening. Colonoscopy is most useful for chronic bleeding or in stable patients when bleeding has ceased. The presence of active bleeding requires several hours of preparation and an experienced colonoscopist. Accuracy of diagnosis is variable but this approach allows the possibility of endoscopic therapy with laser, bipolar electrocoagulation or heater probe.

Abdominal scintigraphy using technetium-99m sulphur colloid or technetium-99m labelled red cells may be helpful to determine fresh bleeding but accurate localization of bleeding is not possible.

Selective mesenteric angiography is the investigation of choice for localization of active bleeding unless the urgency of the situation precludes any further delay. Small intestinal localization of angiodysplasia in the elderly has been useful using this method.

Barium studies can usefully identify small bowel tumours but may delay diagnosis and treatment in the presence of massive bleeding.
Brearley et al.²² have shown that laparotomy is an important diagnostic tool in the management of intestinal bleeding of obscure origin particularly when bleeding is rapid and urgent intervention is indicated. Interoperative endoscopy has been employed to locate small bowel lesions which are impalpable, particularly angiodyplasias. The colonoscope may be passed orally²³ or rectally,²⁴ manipulating the endoscope into the small bowel. Smaller diameter endoscopes can be introduced via an enterotomy, as in the case described above, for more rapid intraluminal bowel inspection or transillumination of the bowel wall to identify vascular abnormalities.²⁵,²⁶

The condition of our patient merited urgent surgery despite the lack of an accurate localization of bleeding preoperatively. Preoperative ileal endoscopy using the gastroscope enabled the involved area of bowel to be visualized and allowed an effective resection.

An unusual acute complication of small bowel lymphoid hyperplasia in a patient with adult Gaucher's disease is presented. Fortunately, the outcome for the patient was successful due to peroperative endoscopic localization of the affected ileum.

References


