Spontaneous rupture of a splenotic nodule

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Summary: A case is presented of spontaneous rupture of splenic tissue occurring 14 years after a splenectomy was carried out for trauma. Spontaneous rupture of a splenotic nodule has not previously been described and it may be added to the list of causes of spontaneous haemoperitoneum. The incidence and function of residual splenic tissue are briefly discussed and other causes of splenic rupture are outlined.

Introduction

Autotransplantation following splenic rupture was described and named splenosis by Buchbinder and Liphoff in 1939 and had been found in dogs as early as 1883. There is some dispute about the incidence of such tissue but several authors have noted that patients who have had a splenectomy for trauma have a lower rate of subsequent serious infection than those for whom this operation has been done for other reasons. Spontaneous rupture of the spleen is a rare, although well-described, event, which may have many causes and which may give rise to diagnostic problems. Although there is one case report of traumatic rupture of a splenotic nodule our case appears to be the only one on record of its spontaneous rupture.

Case report

A 27 year old woman was admitted with a short history of left iliac fossa pain which radiated to her back. She had been taking the oral contraceptive pill and denied the possibility of pregnancy. There was no obvious precipitating cause for the pain and she had been completely well until its onset. Fourteen years prior to this admission she had had a splenectomy for trauma. There was nil else of note in her history. On examination she was pale with a pulse rate of 120 per minute and a blood pressure of 90/60 mmHg. Abdominal examination showed left sided tenderness and guarding. Rectal and vaginal examinations were normal. Her haemoglobin was 10.2 g/dl and her white cell count was $8.1 \times 10^9/l$. The clinical impression was of a ruptured ectopic pregnancy. The patient was resuscitated and taken to theatre. A Pfannenstiel incision was made, 1400 ml of blood were evacuated from the peritoneal cavity and it was determined that the reproductive organs were normal. A left upper paramedian incision was then made. A retroperitoneal haematoma was found in the left upper quadrant and the left colon was mobilized. A splenotic nodule measuring 5 cm in diameter was found in the centre of the haematoma. Its capsule had ruptured and it was excised. A drain was inserted and the wound closed. The patient made an excellent post-operative recovery complicated only by a mild allergic reaction to amoxycillin. This consisted of a skin rash which resolved on withdrawal of the drug. The patient’s white cell count was not elevated at any stage and no specific investigations were carried out. Histology confirmed that the abdominal mass consisted of splenic tissue.

One year later the patient required repair of a small incisional hernia. She has been well since.

Discussion

Livingstone et al. found an incidence of splenosis of 26% in a study of 23 patients who had had a splenectomy for trauma, but higher figures have been quoted by other authors; a scintigraphic study by Solheim and Nerdrum found evidence of it in 11/12 patients splenectomized for trauma. Certainly the incidence and function of splenosis are not sufficient to guarantee protection from overwhelming post-splenectomy sepsis and vaccination or antibiotic treatment is still warranted. Accessory spleens have been found in 20% of patients undergoing splenectomy for staging of Hodgkin’s disease and may give rise to recurrence of those haematological diseases for which the procedure is still carried out.

Spontaneous rupture of the normal spleen is an exceedingly rare event and, as in our patient, has caused diagnostic difficulties in women of

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childbearing years. Spontaneous rupture of the diseased spleen has been described in a variety of conditions and is termed 'pathological rupture'. This often fatal complication has been seen in such generally benign conditions as infectious mononucleosis, chicken pox and salmonella infection as well as such disorders as rheumatoid arthritis, congenital afibrinogenaemia, acquired immune deficiency syndrome, amyloidosis and infective endocarditis. Other disorders which have been reported to number this event among their rare complications include: chronic pancreatitis, hepatitis A and portal hypertension. We are uncertain as to the cause of the rupture in our patient. The clinical course (rapid onset of and recovery from her acute abdomen with an absence of other symptoms) does not suggest any particular aetiological factor. Her white cell count was not elevated during her admission and the differential count was normal. Her reaction to amoxyillin is suggestive of infectious mononucleosis although, as no specific serological investigations were carried out, this putative diagnosis cannot be confirmed.

The findings in our patient raise a number of points. Firstly, as spleenic tissue may regrow, the need for complete removal of the intact spleen (with a thorough check being made for accessory spleens) in cases of those blood disorders for which it is indicated is emphasized. Secondly, although residual spleenic tissue may have some degree of function and the placing of slivers of spleen in the abdomen after splenectomy is worthwhile, it does not offer complete protection and administration of polyvalent pneumococcal vaccine is still recommended. In addition to the mimicking of ruptured ectopic pregnancy by splenic tissue, splenosis can cause further problems for surgeons and gynaecologists by being difficult to distinguish from metastatic carcinoma or endometriosis at laparotomy.

No other case of spontaneous rupture of recrudescent or residual spleenic tissue has been found on record and, of course, ruptured ectopic pregnancy is still the most likely cause of spontaneous haemoperitoneum in women of childbearing age.

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References


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