Spontaneous Appendicocutaneous Fistula

A Case Report

Dr O. M. Tokode* MB, BS
and
Dr O. A. Awojobi+ FMCS (Nig)

ABSTRACT

Ruptured appendicitis is not a common cause of spontaneous enterocutaneous fistula. A case of ruptured retrocaecal appendicitis presenting as an enterocutaneous fistula in a Nigerian woman is presented. The literature on this disorder is also reviewed.

INTRODUCTION

Spontaneous appendicocutaneous fistula following acute appendicitis is a rare clinical entity. It is defined as the primary perforation of the appendix to an adjacent area of the skin, excluding fistulae arising as a sequela of surgically treated appendicitis. However, it has been reported with granulomatous appendicitis, Crohn's disease and when a malignancy in the appendix caused a mucocele, which ruptured into the retroperitoneal space eventually draining via cutaneous sinuses. Enterocutaneous fistula following retrocaecal appendicitis usually involves the caecum or ascending colon.

In the present communication, we report a case of spontaneous appendicocutaneous fistula complicating retrocaecal retroperitoneal appendicitis in a Nigerian woman.

CASE PROFILE

BS, a 38-year-old female farmer presented at Awojobi Clinic Eruwa on 7th September 2000 with passage of undigested food particles through a sore in the right flank of six years duration. The discharge was preceded by a painful right lower quadrant abdominal swelling which spontaneously discharged a lot of pus through the sore. Examination showed a healthy looking woman with a fistula in the right flank just superior to the iliac crest, discharging intestinal contents (Fig 1).

Fistula opening just superior to the iliac crest.

Her packed cell volume was 40% and white cell count of 3000/mm³ with normal differentials. Fistulography using 60% urograftin showed a single tract leading into the proximal ascending colon and the caecum (Fig 2).

Fistulogram showing the tract to the ascending colon and caecum.

Exploratory laparotomy revealed a retroperitoneal retrocaecal appendix with a colocutaneous fistula.
tract into the ascending colon at the tip of the shrunken appendix which was adherent to the colon. No intraperitoneal pathology was found. The defect in the colon was closed in two layers with 0 silk and appendicectomy was performed. She made uneventful recovery and was discharged on the seventh postoperative day.

Histopathology report read: Piece of appendicectomy specimen measuring 5.5cm in length and 1.3cm in its widest width. The serosal surface is moderately congested and the lumen in the middle third is distended with faecal materials. Sections show moderate infiltration of appendiceal mucosa and submucosa by plasma cells and eosinophils, polymorphs, with mild involvement of the muscularis propria. There is mucosal lymphoid hyperplasia with prominent germinal centre. These features are consistent with eosinophilic appendicitis.

She has been followed up for four years and her condition has remained satisfactory with no recurrence of the fistula.

DISCUSSION
An intestinal fistula is an abnormal tract that communicates between the intestinal mucosa and another epithelial surface. In enterocutaneous fistula the other epithelial surface is the skin and intestinal contents are discharged externally.

The common acquired causes of enterocutaneous fistula include strangulated groin hernia especially femoral hernia, tuberculosis and other granulomatous infections, diverticular disease of the bowel, Crohn’s disease, carcinoid tumour and carcinoma of the caecum and appendix.10,15

Appendiceal fistula has been the presentation of pseudomyxoma retroperitonei a late complication of mucinous adenocarcinoma of the appendix.11,12

Most enterocutaneous fistulas in this environment have been complications of surgery for peritonitis due to typhoid ileitis, perforated appendicitis, perforated duodenal ulcer, penetrating injuries of the abdomen, septic criminal abortion and intestinal amoebiasis.13,16,21 It has followed incomplete appendicectomy.8

Spontaneous appendicocutaneous fistula complicating acute appendicitis is rare. The cutaneous opening has been in the buttock,4 right iliac fossa,4 the right groin6 and in the right flank as in our case report. More common appendiceal fistulas have involved the urinary bladder, colon, duodenum and ileum.6

The effect of the fistula on the patient is least pronounced being a low output fistula the loss of fluids, electrolytes and nutrients is minimal.

Diagnosis is confirmed by fistulography. This displays the fistula tract and its connection to the gut, its level and the presence of associated abscess cavity or distal obstruction. Barium enema with air contrast helps to detect any associated colonic disease like diverticulitis, Crohn’s disease or polyposis coli.

The treatment of appendiceal fistula is appendicectomy and closure of the colonic defect. No recurrence is expected if there is no other associated pathology.

REFERENCES
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