Chronic Intestinal Idiopathic Pseudo-obstruction (CIIPO) Complicated by Sigmoid Volvulus

Christos Konstantinidis*

Surgical Department, Hippocratio General Hospital of Athens, 114 Vasilissis Sofias str, Athens, Region of Attica, Greece

*Corresponding author: drchriskons@yahoo.gr

Abstract A 17-year-old male patient with chronic intestinal idiopathic pseudo-obstruction was admitted to the emergency department clinically deteriorated due to sigmoid volvulus. Initial resuscitative efforts followed by endoscopy failed to detorse and decompress the bowel. At the followed emergency laparotomy the volvulus was derotated and a Hartmann’s procedure with left hemicolectomy was performed. The patient after 3 months underwent stoma closure. At one-year follow-up the patient showed clinical improvement.

Keywords: constipation, megacolon, pseudo obstruction, sigmoid volvulus


1. Introduction

Intestinal pseudo-obstruction (megacolon) refers to bowel dilation without known mechanical occlusion of the lumen. It can be acute (AIPO), as in Ogilvie’s syndrome or chronic (CIPO), either primary or secondary (electrolyte disorders, viruses, toxic causes etc). Primary CIPO, can be congenital or acquired, due to known histopathological abnormalities: neuropathies (intrinsic and extrinsic gastrointestinal nerve pathways affected), mesenchymopathies (interstitial cells of Cajal affected) and myopathies (smooth muscle cells affected) [1].

Chronic intestinal idiopathic pseudo-obstruction (CIIPO) describes permanently distended bowel of unknown origin. In most cases CIIPO is sporadic, while familial forms (autosomal dominant, autosomal recessive and X-linked transmission) have also been described [2,3].

Herein, it is presented a case of sigmoid volvulus in an adult with known CIIPO.

2. Case Presentation

A 17-year-old male patient presented to the emergency department with marked abdominal distention, vomiting and referred faecal incontinence for several days. From his medical history he had had mental retardation and chronic constipation since he was one year old. He was receiving long-term laxatives for the treatment of constipation. He has been diagnosed as having non-Hirschsprung congenital megacolon, after mentioned thorough investigation in the past.

On admission his general clinical condition was bad. The patient was lethargic, dehydrated with low blood pressure and tachycardia, tachypneic but afebrile. On clinical examination the abdomen was diffusely distended and tympanic on percussion, with no sign of peritonitis.

Plain abdominal radiograph (Figure 1) showed grossly dilated intestinal loops. Subsequent abdominal scan (Figure 2, Figure 3) indicated sigmoid volvulus. Initially, the patient was resuscitated with intravenous fluids while gastric decompression attempted via nasogastric tube. Flexible sigmoidoscopy in the aim of intestinal detorsion and decompression via was unsuccessful. Due to patient’s clinical deterioration, he underwent emergency laparotomy, at which massive dilation of bowel loops were seen (Figure 4). There were no signs of bowel ischemia or
perforation. The sigmoid colon was derotated and left hemicolecctiony through Hartmann’s procedure was performed. Also, an accidental splenectomy was occurred. Postoperative period was uneventful and the patient was discharged 2 weeks after admission. He was referred to a specialist center for cytogenetic (chromosomal) analysis, which didn’t reveal morphological anomalies of autosomal or X-linked chromosomes. Histopathological examination of the resected specimen was not indicative of Hirschsprung’s disease.

Figure 2. Abdominal scan indicating sigmoid volvulus

Figure 3. Another abdominal scan view indicating sigmoid volvulus
Stoma closure was performed 3 months postoperatively. At one-year follow-up the patient is in good condition, with gained weight and less requirements for laxatives.

3. Discussion

Among large bowel, sigmoid colon is more often affected from volvulus and is more frequent in adults [4]. Initial colonoscopic attempt for decompression and detorsion of the affected bowel segment is the treatment of choice, when it is possible and the clinical situation of the patient allows it. In suspicious or obvious peritonitis, as well as when the patient deteriorates, urgent laparotomy is indicated [5,6]. While Hirschsprung’s disease represents the most known form of congenital megacolon, affecting from distal part of sigmoid colon and rectum to total large bowel [7], chronic intestinal pseudo-obstruction (CIPO) is a rare syndrome, with a particular subgroup called chronic idiopathic intestinal pseudo-obstruction (CIPO) [8]. The latter characterizes chronic constipation accompanied with persistent bowel dilation in the absence of known cause [9]. It is diagnosed by exclusion of others relevant modalities [10]. The main treatment option is conservative, with long-term laxatives. In urgent situations, such as in the prescribed case, colonoscopic decompression followed, in failed attempts, by surgical intervention. Surgical options include stoma formation or colectomy with anastomosis.

References