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CASE REPORT

Pneumatosis coli causing pneumoperitoneum

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Abstract

This is a case of a 54-year-old gentleman who presented to an outside hospital emergency department with lower abdominal pain. Computed tomography imaging showed a small amount of intraperitoneal free air and cystic pneumatosis coli. He was admitted, managed conservatively with intravenous antibiotics, and then discharged home after his symptoms improved. Elective laparoscopic sigmoid colectomy was subsequently performed with intraoperative findings of partial sigmoid volvulus and extensive pneumatosis coli of the sigmoid colon. Pneumoperitoneum was determined to be from ruptured intramural cysts. The etiology of pneumatosis coli was likely from chronic sigmoid volvulus.

INTRODUCTION

Pneumatosis cystoides intestinalis (PCI) is a rare finding characterized by multiple air-filled cysts in the submucosa, subserosa and/or muscularis propria of the intestine. Recent studies report that PCI or pneumatosis intestinalis is more frequently found in the colon than the small bowel [1, 2]. When located in the colon, it is specifically termed pneumatosis coli.

PCI is not well understood. Symptoms associated with it are non-specific, commonly including abdominal pain and distension, diarrhea, bloody stool and constipation [1]. The underlying pathology is diverse and not always identified. The presence of PCI can be an incidental finding of no clinical significance; however, it can also be a sign of an underlying disease process such as bowel necrosis, mesenteric ischemia and bowel obstruction which can be life-threatening. When a serious underlying intra-abdominal pathology is suspected, urgent surgical intervention is warranted.

We present a case of pneumatosis coli and pneumoperitoneum in an otherwise healthy male patient.

CASE REPORT

A 54-year-old gentleman with 30 pack year history of smoking, chronic constipation and symptoms of irritable bowel syndrome presented to an outside hospital emergency department (ED) with 5 days of lower abdominal pain and cramping. The pain was initially very mild in nature, but suddenly became severe a few days after onset. The pain then subsided, but did not completely resolve.

In the ED, heart rate was normal. He was afebrile with normal white blood cell count and C-reactive protein level. On examination, there was no evidence of peritonitis. Computed tomography (CT) of the abdomen pelvis demonstrated findings concerning for colonic perforation; there was a small amount of free intraperitoneal air scattered throughout the abdomen and pelvis and prominent cystic pneumatosis coli of the sigmoid colon (Fig. 1). The etiology was indeterminate. He was admitted and managed conservatively with intravenous (IV) antibiotics. His symptoms improved, and he was discharged home on hospital Day 4.

He presented to our clinic 2 days after discharge for further work-up complaining of persistent mild symptoms of abdominal pain. Flexible sigmoidoscopy was performed showing approximately a 7 cm segment of sigmoid colon with numerous polypoid appearing lesions with grossly normal appearing overlying mucosa (Fig. 2). These lesions were biopsied with pathology revealing hyperplastic changes without dysplasia.
Notably, his last colonoscopy was approximately 10 months ago which demonstrated tubular adenomas, one in the descending colon and the other in the rectum.

Notably, his last colonoscopy was approximately 10 months ago which demonstrated tubular adenomas, one in the descending colon and the other in the rectum.
Given his symptoms persisted, he was taken to the operating room electively for additional evaluation and potential therapeutic intervention. Intraoperatively, the sigmoid colon was notably redundant with a narrow mesenteric stalk. There were findings of chronic non-obstructing sigmoid volvulus with a twisted and inflamed sigmoid mesentry. No evidence of gross perforation was identified. Extensive PCI of the sigmoid colon was apparent externally and upon opening of the specimen on the back table (Fig. 3). Laparoscopic sigmoid colectomy with end-to-end colorectal anastomosis was performed.

The specimen was sent to pathology. Air pockets in the submucosa ranging from 0.3 to 1.5 cm in diameter were identified (Fig. 4). Microscopy revealed multiple intramural empty cysts lined by histiocytes and multinucleated giant cells, consistent with PCI (Figs 5 and 6). There was no evidence of gross perforation identified on final pathology.

**DISCUSSION**

In this case report, we describe a patient who initially presented with symptomatic pneumatosis coli and pneumoperitoneum of undetermined etiology. Despite the imaging findings, his entire clinical picture did not warrant emergent surgical intervention, and he was managed conservatively as an inpatient. He was discharged home and additional work-up was performed on an outpatient and elective basis, revealing multiple sigmoid hyperplastic polyps without evidence of transmural colonic perforation.

Pneumatosis coli in association with PCI can be secondary to viscus perforation or ruptured intramural cysts [3]. When this patient initially presented to the ED, he had free air on CT concerning for colonic perforation. However, further work-up determined the source of pneumoperitoneum was likely from ruptured intramural cysts as there was no evidence of gross transmural perforation intraoperatively and on final surgical pathology of the sigmoid colon. This correlates with his benign clinical picture.

The underlying pathology of PCI is diverse and not always identified. Sigmoid volvulus has been reported as both an etiology and a complication of PCI [4]. Here, we favor chronic sigmoid volvulus as the etiology rather than a complication of PCI. He had a redundant sigmoid colon, short mesenteric attachment and history of constipation, predisposing him to sigmoid volvulus. His presentation of recurrent abdominal pain with resolution may be due to recurrent torsion and spontaneous detorsion.

This case describes chronic sigmoid volvulus as a potential etiology of PCI and ruptured intramural cysts resulting in pneumoperitoneum.

**REFERENCES**