Pneumatosis Intestinalis as the Initial Presentation of Systemic Sclerosis: A Case Report and Review of the Literature

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1. Introduction

Pneumatosis intestinalis (PI) is an uncommon pathology, first described in 1730 [1], characterized by the presence of gas within the intestinal wall, usually in the mucosa and submucosal layers of the antimesenteric border [2]. Primary PI is a benign idiopathic condition [3], while secondary PI can be associated with various underlying conditions, including necrotising enterocolitis, obstructive lung disease, inflammatory bowel disease, and connective tissue diseases [4-6]. Even though the association between PI and connective tissue diseases is well known [2, 7], antiphospholipid syndrome is the only mixed connective tissue disorder in which, it has been reported as the initial presentation [8]. To our knowledge, this is the first report of PI being the initial presentation of systemic sclerosis.

2. Case Presentation

The patient was a 75-year-old female with an 8-month history of worsening dysphagia, to both solids and liquids and epigastric pain, as well as other nonspecific symptoms. Initial investigations with an oesophagogastroduodenoscopy diagnosed Candida oesophagitis and also identified an extrinsic compression of the gastric antrum. Subsequently a CT scan of the abdomen and pelvis showed moderately dilated small bowel loops and PI. Due to the patient’s stability, non-critical clinical condition, conservative management was instituted. More detailed investigations confirmed the diagnosis of systemic sclerosis with positive anticentromeric and antinuclear antibodies. The patient improved on methotrexate and was discharged with appropriate outpatient follow-up. Discussion. PI is a rare but well-documented pathology associated with connective tissue diseases, such as systemic sclerosis. In most cases, conservative management is preferable to surgical intervention, depending on the patient’s clinical presentation and progress. This is the first report of PI being the initial presentation of a patient with systemic sclerosis responsive to conservative management.
loops with gas within the intestinal wall (Figure 1). Since performed. The CT showed moderately dilated small bowel extrinsic compression of the gastric antrum and therefore a gastric mucosa on biopsy. However, it also identified an COPD) and mild chronic non-active inflammation of the oesophagitis (possibly due to the inhaled steroids for the patient. The patient was initially investigated with an oesophagogastroduodenoscopy (OGD), which diagnosed Candida oesophagitis (possibly due to the inhaled steroids for the COPD) and mild chronic non-active inflammation of the gastric mucosa on biopsy. However, it also identified an extrinsic compression of the gastric antrum and therefore a computer tomography (CT) of the abdomen and pelvis was performed. The CT showed moderately dilated small bowel loops with gas within the intestinal wall (Figure 1). Since the patient’s clinical condition was stable and did not demonstrate signs of peritonism, a decision for conservative management was made. After excluding life-threatening and acute causes of PI, such as infection or ischemia, further investigations were performed guided towards the more uncommon pathologies. A screen for autoimmune diseases showed positive results for the anticientromeric and antinuclear antibodies and the diagnosis of systemic sclerosis was made. With the diagnosis confirmed, a more detailed medical history was obtained, specifically guided to reveal any symptoms of systemic sclerosis, with no success. Similarly, no signs related to the disease, such as skin thickening, calcinosis, or telangectasia, were identified. The patient improved on methotrexate and was discharged after a total of 4 weeks hospital stay with appropriate outpatient follow-up.

3. Discussion

Pneumatosis intestinalis (PI) is an uncommon pathology characterised by the presence of gas within the intestinal wall, associated with a variety of clinical disorders [2, 4, 5], including connective tissue diseases [6]. Among the latter, PI has been described in progressive systemic sclerosis, also reported as a late complication and a poor prognostic indicator [8]; systemic lupus erythematosus, where it is suggested to reflect ischemic necrosis of the bowel due to vasculitis [9]; mixed connective tissue disease [2, 7, 9–11], where it may be associated with a better prognosis [12, 13], while it is often associated with corticosteroid administration [14].

Although the pathophysiology of PI is uncertain, two main theories have been proposed. The mechanical theory advocates that increased luminal pressure caused by intestinal obstruction allows gas to penetrate into the submucosal space through a mucosal breach [15]. The bacterial theory supports that gas-producing bacteria, such as Clostridium difficile or Clostridium perfringens, invade the submucosal layer through mucosal rents and produce gas within the intestinal wall [16]. Finally, it has been suggested that long-term administration of corticosteroids possibly induces atrophy of the intestinal mucosa, sometimes resulting in a mucosal defect and the subsequent translocation of gas into the submucosal layer [17, 18]. This could be the explanation also in our case, where the patient was on long-term use of corticosteroids for her COPD.

The clinical importance of PI is that it may be misdiagnosed or, if identified correctly, its significance may be misinterpreted, leading to unnecessary surgical intervention [19]. Many cases of colonic gas cysts present with symptoms of large bowel dysfunction, (usually diarrhea, colicky abdominal pain, and excessive flatulence) and, occasionally, rectal bleeding. Resection of the affected segment of the colon usually, but not always, relieves the symptoms, but in many cases spontaneous relief of symptoms can also occur [20, 21]. On the contrary, PI can also be found in cases where the viability of the bowel is threatened [6]. In these cases, it is considered to be a manifestation of a form of intestinal gas gangrene [22], and it usually indicates the need of emergency laparotomy, depending on the patient’s clinical signs. In our case, the patient did not exhibit any signs of peritonism or bowel ischaemia and therefore conservative management was preferred.

In conclusion, PI is a rare but well-documented pathology associated with connective tissue diseases, such as systemic sclerosis. In most cases, conservative management is preferable to surgical intervention, however, this decision depends on the patient’s clinical presentation and progress. This is the first report of PI being the initial presentation of a patient with systemic sclerosis responsive to conservative management.

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